



US008522832B2

(12) **United States Patent**
Lopez et al.

(10) **Patent No.:** **US 8,522,832 B2**
(45) **Date of Patent:** **Sep. 3, 2013**

(54) **FLUID TRANSFER DEVICES AND METHODS OF USE**

(75) Inventors: **George A. Lopez**, Laguna Beach, CA (US); **Thomas F. Fangrow, Jr.**, Mission Viejo, CA (US); **Peter Leissling**, Schalksmuehle (DE); **Matthias Janssen**, Meinerzhagen (DE)

(73) Assignee: **ICU Medical, Inc.**, San Clemente, CA (US)

(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 543 days.

(21) Appl. No.: **12/845,548**

(22) Filed: **Jul. 28, 2010**

(65) **Prior Publication Data**

US 2011/0062703 A1 Mar. 17, 2011

Related U.S. Application Data

(60) Provisional application No. 61/229,701, filed on Jul. 29, 2009, provisional application No. 61/354,648, filed on Jun. 14, 2010.

(51) **Int. Cl.**
B65B 1/04 (2006.01)
A61B 19/00 (2006.01)

(52) **U.S. Cl.**
USPC **141/9**; 141/27; 141/94; 141/329;
604/416; 604/905

(58) **Field of Classification Search**
USPC 141/9, 23–27, 329–330, 301–302,
141/94, 98, 318–322; 604/408, 410–411,
604/407, 416, 905

See application file for complete search history.

(56) **References Cited**

U.S. PATENT DOCUMENTS

1,923,501 A 8/1933 Perry
4,005,710 A 2/1977 Zeddies et al.

(Continued)

FOREIGN PATENT DOCUMENTS

DE 20 2004 014 868 11/2004
EP 1 997 471 12/2008

(Continued)

OTHER PUBLICATIONS

International Search Report and Written Opinion re PCT Application No. PCT/US2010/043451, mailed Nov. 30, 2011.

(Continued)

Primary Examiner — Timothy L. Maust

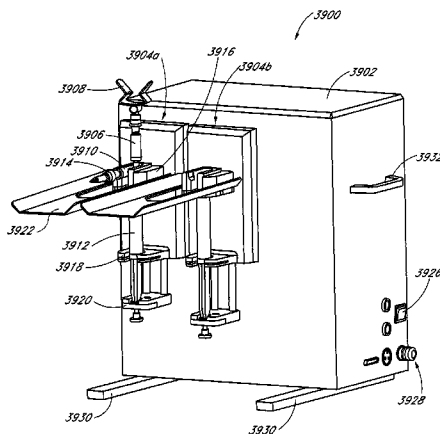
Assistant Examiner — Timothy P. Kelly

(74) *Attorney, Agent, or Firm* — Knobbe, Martens, Olson & Bear, LLP

(57) **ABSTRACT**

Some embodiments disclosed herein related to a device for transferring precise amounts of fluid from at least one source container to at least one target container. In some embodiments, the fluid is first transferred from the source container (e.g., a vial) through a connector to an intermediate measuring container (e.g., a syringe). In some embodiments air can pass through an air inlet and enter the vial to compensate for the volume of fluid withdrawn from the vial. An air check valve or a bag or a filter can prevent the fluid from escaping through the air inlet. The precisely measured amount of fluid can then be transferred from the intermediate measuring container to the target container (e.g., an IV bag). In some embodiments the connector can include a source check valve and a target check valve to direct fluid first from the source container to the intermediate measuring container and then from the intermediate measuring container to the target container. Some embodiments of the device can include a motor and a controller for automatically actuating a plunger of the syringe to transfer the desired amount of fluid.

21 Claims, 107 Drawing Sheets



(56)

References Cited

U.S. PATENT DOCUMENTS

4,084,606 A * 4/1978 Mittleman 137/102
 4,190,048 A 2/1980 Sampson
 4,306,705 A 12/1981 Svensson
 4,559,043 A 12/1985 Whitehouse et al.
 4,561,856 A 12/1985 Cochran
 4,666,429 A 5/1987 Stone
 4,670,007 A 6/1987 Wheeldon et al.
 4,683,916 A 8/1987 Raines
 4,755,172 A 7/1988 Baldwin
 4,778,450 A 10/1988 Kamen
 4,819,684 A 4/1989 Zaugg et al.
 4,863,429 A 9/1989 Baldwin
 4,922,975 A 5/1990 Polaschegg
 4,969,874 A 11/1990 Michel et al.
 4,976,590 A 12/1990 Baldwin
 4,995,268 A 2/1991 Ash et al.
 5,024,347 A 6/1991 Baldwin
 5,037,390 A 8/1991 Raines et al.
 5,114,580 A 5/1992 Ahmad et al.
 5,176,658 A 1/1993 Ranford
 5,256,155 A 10/1993 Yerlikaya et al.
 5,334,211 A 8/1994 Shiber
 5,336,201 A 8/1994 von der Decken
 5,431,201 A 7/1995 Torchia et al.
 5,439,451 A 8/1995 Collinson et al.
 5,609,572 A 3/1997 Lang
 5,676,346 A 10/1997 Leinsing
 5,782,816 A 7/1998 Werschmidt et al.
 5,807,312 A 9/1998 Dzwonkiewicz
 5,897,526 A 4/1999 Vaillancourt
 5,904,666 A 5/1999 DeDecker et al.
 5,947,951 A 9/1999 Ortiz et al.
 5,968,014 A 10/1999 Neftel et al.
 5,989,237 A 11/1999 Fowles et al.
 6,059,747 A 5/2000 Bruggeman et al.
 6,110,153 A 8/2000 Davis et al.
 6,123,685 A 9/2000 Reynolds
 6,132,404 A 10/2000 Lopez
 6,152,900 A 11/2000 Mayer
 6,221,041 B1 4/2001 Russo
 6,245,048 B1 6/2001 Fangrow, Jr. et al.
 6,302,864 B1 10/2001 Nowosielski
 6,425,497 B1 7/2002 Chu et al.
 6,474,375 B2 11/2002 Spero et al.
 6,485,472 B1 11/2002 Richmond
 6,585,229 B2 7/2003 Cote, Sr. et al.
 6,590,167 B2 7/2003 Clare
 6,623,455 B2 9/2003 Small et al.
 6,629,956 B1 10/2003 Polidoro et al.
 6,651,956 B2 11/2003 Miller
 6,663,586 B2 12/2003 Verkaart et al.
 6,689,108 B2 2/2004 Lavi et al.
 6,813,868 B2 11/2004 Baldwin et al.
 6,854,620 B2 * 2/2005 Ramey 222/63
 6,877,530 B2 4/2005 Osborne et al.
 6,908,459 B2 6/2005 Harding et al.
 6,915,823 B2 7/2005 Osborne et al.
 6,948,522 B2 9/2005 Newbrough et al.
 6,953,450 B2 10/2005 Baldwin et al.
 6,991,002 B2 1/2006 Osborne et al.
 6,994,315 B2 2/2006 Ryan et al.
 7,017,623 B2 3/2006 Tribble et al.
 7,117,902 B2 10/2006 Osborne
 7,128,105 B2 10/2006 Tribble et al.
 7,163,035 B2 1/2007 Khan et al.
 7,194,336 B2 3/2007 DiGianfilippo et al.
 7,317,967 B2 1/2008 DiGianfilippo et al.
 7,343,224 B2 3/2008 DiGianfilippo et al.
 7,343,943 B2 3/2008 Khan et al.
 7,351,226 B1 4/2008 Herskowitz
 7,392,638 B2 7/2008 Baldwin et al.
 7,396,051 B2 7/2008 Baldwin et al.
 7,418,981 B2 9/2008 Baker et al.
 7,442,186 B2 10/2008 Blomquist
 7,499,581 B2 3/2009 Tribble et al.

7,538,858 B2 5/2009 Mackey
 7,610,115 B2 10/2009 Rob et al.
 7,681,606 B2 3/2010 Khan et al.
 7,703,483 B2 4/2010 Hartman et al.
 7,789,850 B2 9/2010 Roger
 7,814,731 B2 10/2010 Bender et al.
 7,882,863 B2 * 2/2011 Pestotnik et al. 141/318
 7,900,658 B2 3/2011 Osborne et al.
 7,913,720 B2 3/2011 Tribble et al.
 7,967,202 B2 6/2011 Durrell et al.
 7,997,304 B2 8/2011 Ranalletta et al.
 8,091,860 B2 * 1/2012 Thompson et al. 251/129.04
 8,140,351 B2 3/2012 Tribble et al.
 8,151,835 B2 4/2012 Khan et al.
 2002/0179544 A1 12/2002 Johnson et al.
 2004/0035743 A1 2/2004 Tighe et al.
 2004/0116891 A1 6/2004 Curutcharry
 2005/0059952 A1 3/2005 Giuliano et al.
 2005/0096627 A1 * 5/2005 Howard 604/500
 2005/0131357 A1 6/2005 Denton et al.
 2005/0252574 A1 11/2005 Khan et al.
 2006/0064053 A1 3/2006 Bollish et al.
 2006/0259195 A1 11/2006 Eliuk et al.
 2007/0088252 A1 4/2007 Pestotnik et al.
 2007/0106244 A1 5/2007 Mosler et al.
 2008/0065006 A1 3/2008 Roger et al.
 2008/0114328 A1 5/2008 Doherty et al.
 2008/0125897 A1 5/2008 DiGianfilippo et al.
 2008/0169044 A1 7/2008 Osborne et al.
 2008/0195416 A1 8/2008 Tribble et al.
 2008/0199353 A1 8/2008 Mlodzinski et al.
 2008/0287920 A1 11/2008 Fangrow et al.
 2009/0012449 A1 1/2009 Lee et al.
 2009/0050216 A1 2/2009 Trocki et al.
 2009/0067973 A1 3/2009 Eliuk et al.
 2009/0069743 A1 3/2009 Krishnamoorthy et al.
 2009/0082649 A1 3/2009 Muller et al.
 2009/0088687 A1 4/2009 Yardimci et al.
 2009/0099547 A1 4/2009 Radmer
 2009/0101576 A1 4/2009 Rohde et al.
 2009/0126825 A1 5/2009 Eliuk et al.
 2009/0145509 A1 6/2009 Baker et al.
 2009/0149743 A1 * 6/2009 Barron et al. 600/431
 2009/0163860 A1 6/2009 Patrick et al.
 2009/0177149 A1 7/2009 Childers et al.
 2009/0198215 A1 8/2009 Chong et al.
 2009/0254031 A1 10/2009 Lee
 2009/0306621 A1 12/2009 Thome, Jr. et al.
 2010/0049157 A1 2/2010 Fangrow
 2010/0245056 A1 9/2010 Braun et al.
 2010/0276034 A1 11/2010 Gonnelli et al.
 2010/0280430 A1 11/2010 Caleffi et al.
 2011/0087164 A1 4/2011 Mosler et al.
 2011/0112501 A1 5/2011 Garfield et al.
 2011/0178493 A1 7/2011 Okiyama
 2011/0196304 A1 8/2011 Kramer et al.
 2012/0078091 A1 3/2012 Suchecki et al.
 2013/0006214 A1 1/2013 Garfield et al.

FOREIGN PATENT DOCUMENTS

WO WO 99/63547 12/1999
 WO WO 2005/123162 12/2005
 WO WO 2007/061424 5/2007
 WO WO 2008/128074 10/2008

OTHER PUBLICATIONS

International Preliminary Report on Patentability and Written Opinion re PCT Application No. PCT/US2010/043451, mailed Feb. 9, 2012.
 User Guide for medOC 1xx Basic, Neo Care Medical Products GmbH, Version 06.2008, 23 pages.
 User Manual for medOC 3xx /6xx /8xx, Neo Care Medical Products GmbH, Version 05.2008, 44 pages.
 Technical Data sheet for Analog Amplifiers Type VA, models V8-C and V8-D, STM Sensors dated Dec. 2007, 4 pages.
 Technical Data sheet for Through Beam Sensors Type G2, 645 nm, STM Sensors dated Sep. 2008, 2 pages.

Technical Data sheet for Through Beam Sensors Type G2, 1480 nm, STM Sensors dated Dec. 2009, 2 pages.

Product detail for "Summit Medical DirectFlow" micro infusion extension set from Summit Medical Technologies, downloaded on May 10, 2010 from <http://summitmedtech.com/p6line.php>, 1 page.

Product detail for "NAMIC® Closed Fluid Systems" from Navilyst Medical, downloaded on May 11, 2010 from <http://www.navilystmedical.com/Products/index.cfm/19>, 2 pages.

Smith, "Lifesaving Cancer Drugs May Put Workers' Lives at Risk," downloaded on Jul. 12, 2010 from <http://www.msnbc.msn.com/id/38114586/ns/health-cancer>, 7 pages.

Product detail for "RapidFill™ Automated Syringe Filler," from Baxa, downloaded on Mar. 31, 2010 from <http://www.baxa.com/PharmacyProducts/AutomatedFillingSystems/ProductDetail/?id=B1>, 2 pages.

Integra Brochure, from Eurospital, Brochure acquired in Mar. 2012. Autotec 50, from KRZ, Dec. 6, 2007.

Pinnacle TPN Management System, from B Braun, downloaded May 5, 2009 from <http://www.bbraunusa.com/index.cfm?uuid=7386ADF065B05CD0D22AF700339AA4092>, 1 page.

Exacta-Mix 2400, from Baxa, which appears to have a date of 2007, 2 pages.

Riva, downloaded in Apr. 2009 from <http://www.rivasystem.com>, 6 pages.

Fox, Brent I., "Pharmacy Automation and Technology: Automated Intravenous Preparation: Robots for the Pharmacy," Hospital Pharmacy, vol. 44, Mar. 2009, pp. 255-257.

Flickinger, Bruce, "Misperceptions Cloud the Issue of Sterile Drug Compounding," Jun. 2007.

CytoCare, by Health Robotics, Brochure, Date Unknown, downloaded on May 25, 2012 from <http://www.health-robotics.com/smarteredit/downloads/en/cytocare7.pdf>, 6 pages.

International Search Report and Written Opinion dated Feb. 19, 2013, International Application No. PCT/US2012/071493, filed Dec. 21, 2012.

International Search Report and Written Opinion re PCT Application No. PCT/US2012/071493, mailed Feb. 19, 2013.

* cited by examiner

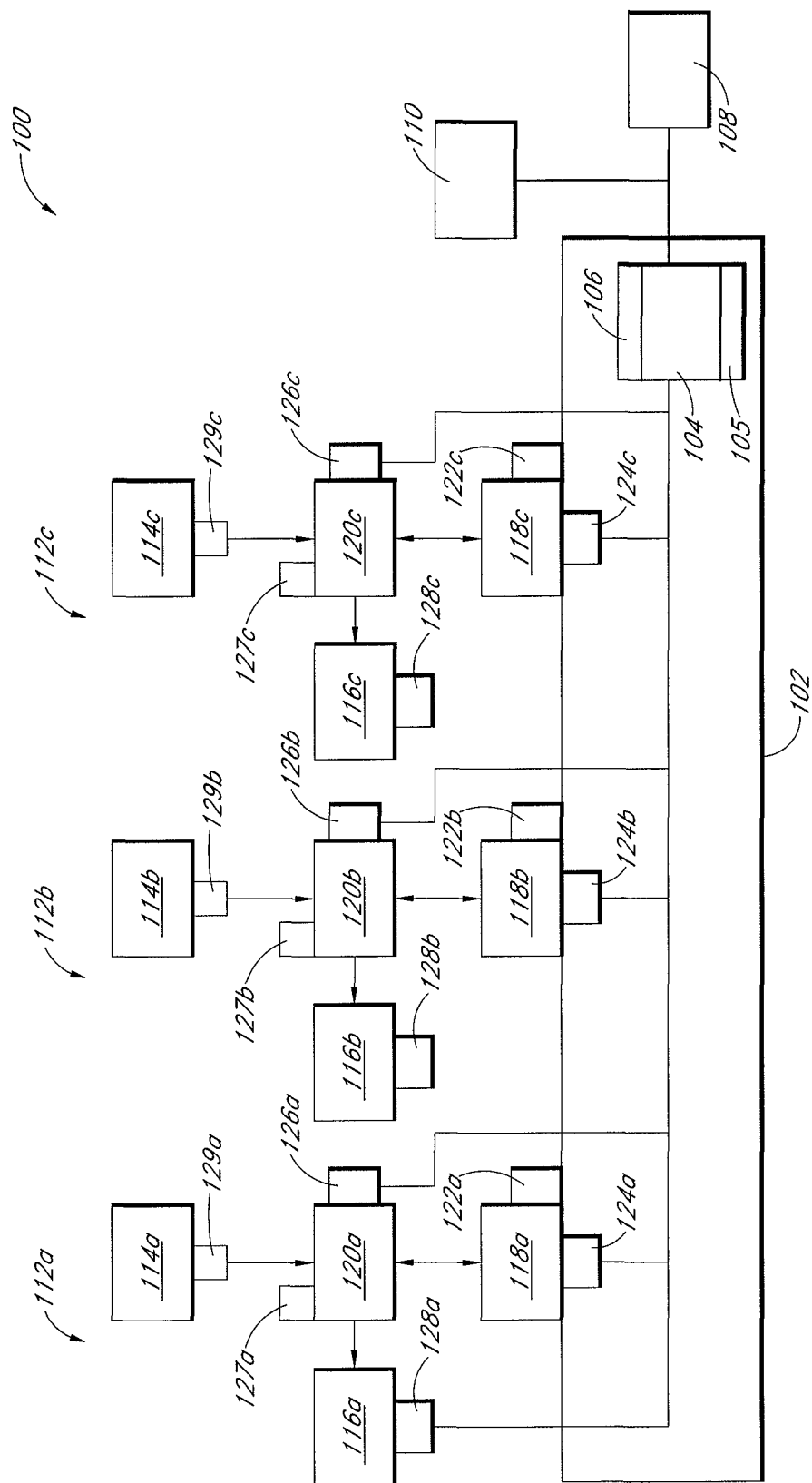


FIG. 1

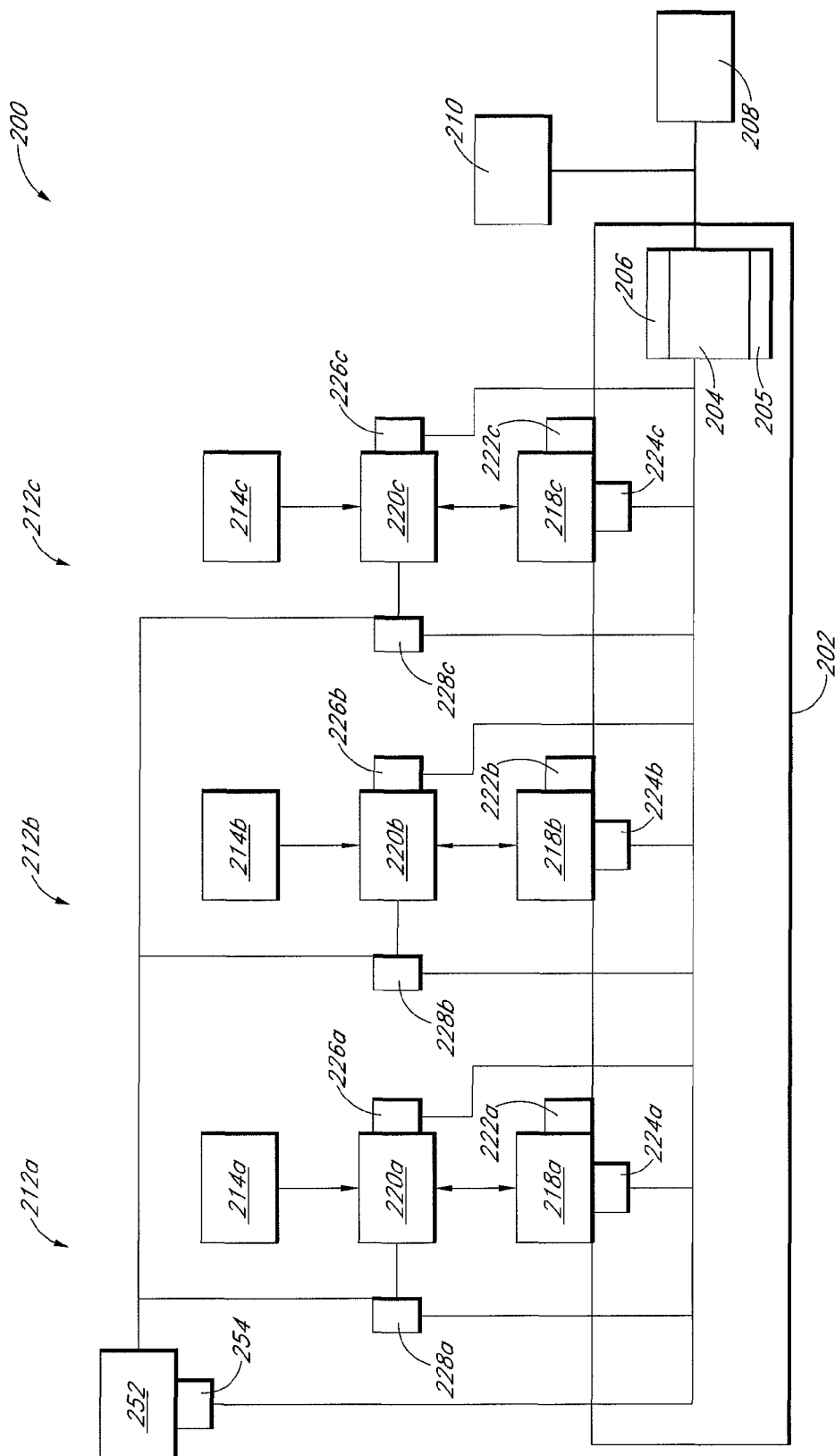


FIG. 2

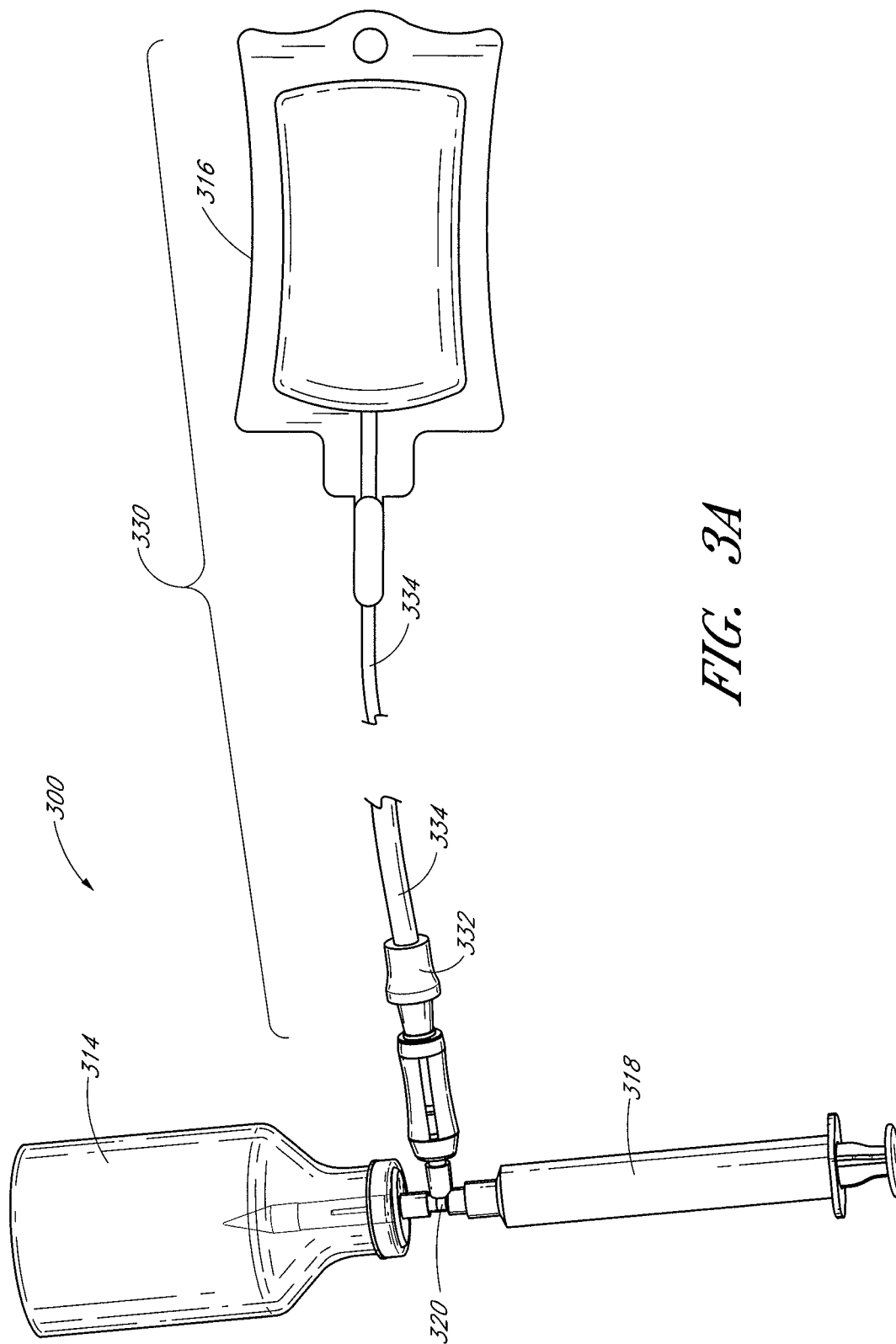


FIG. 3A

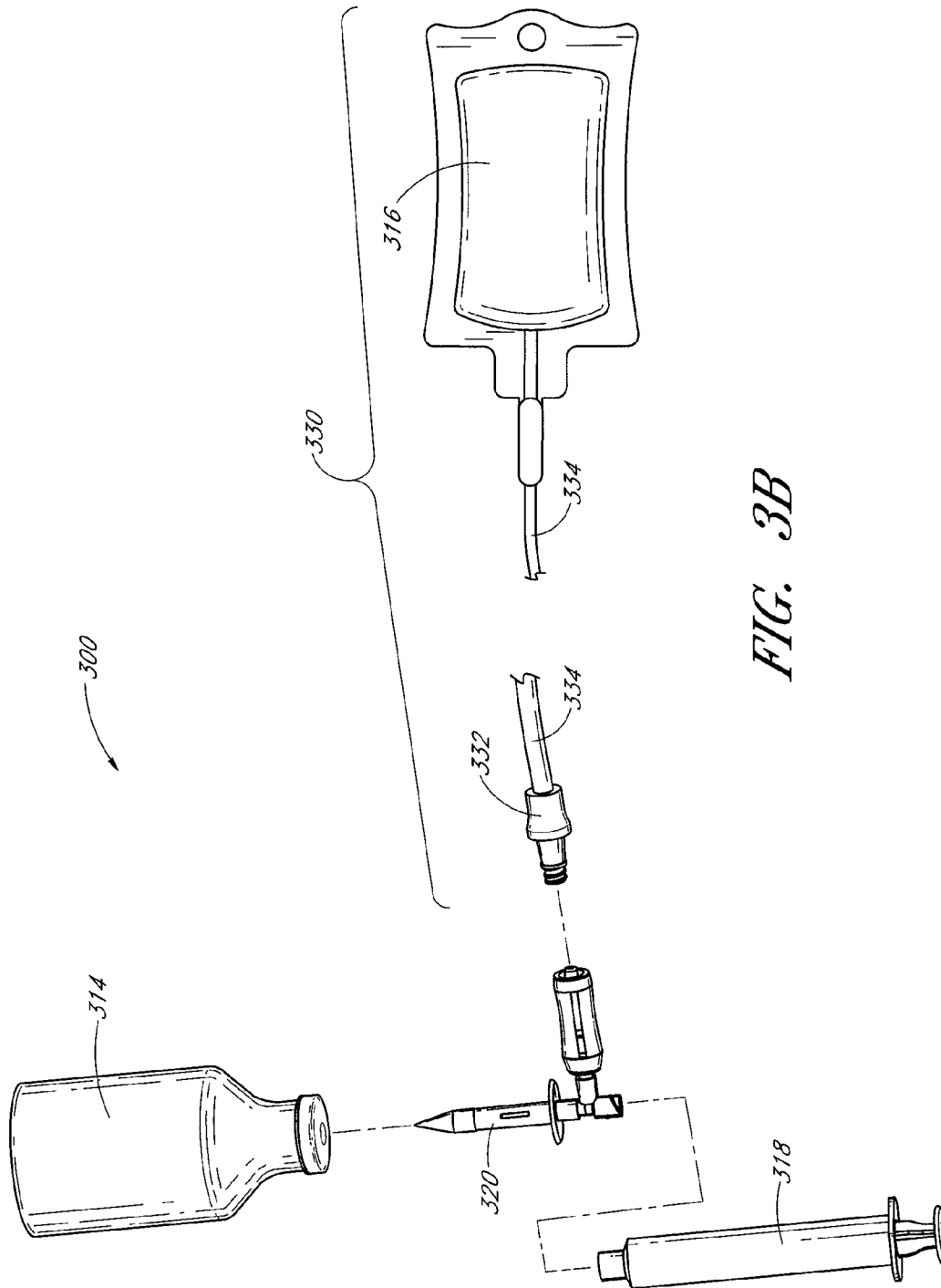
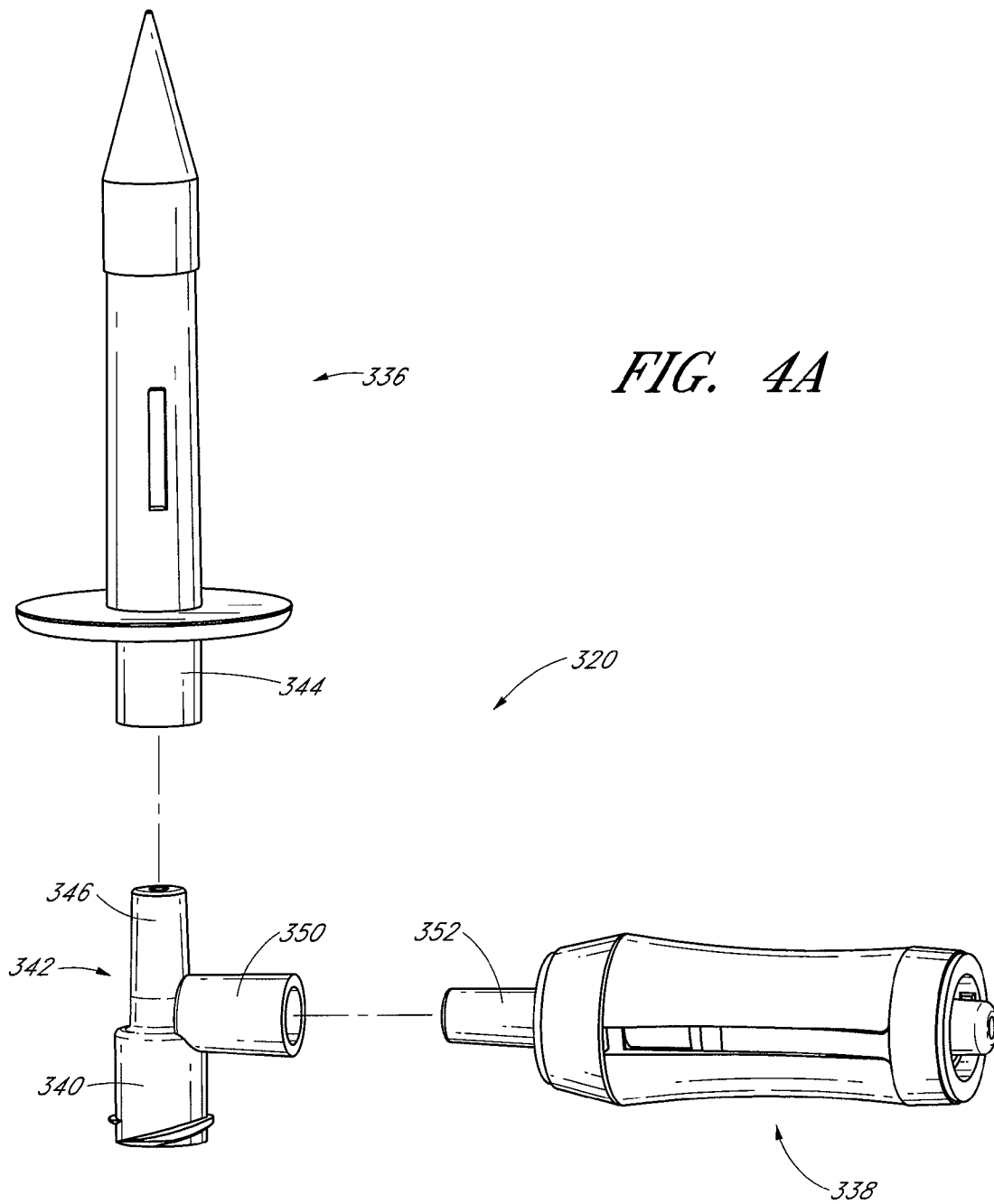
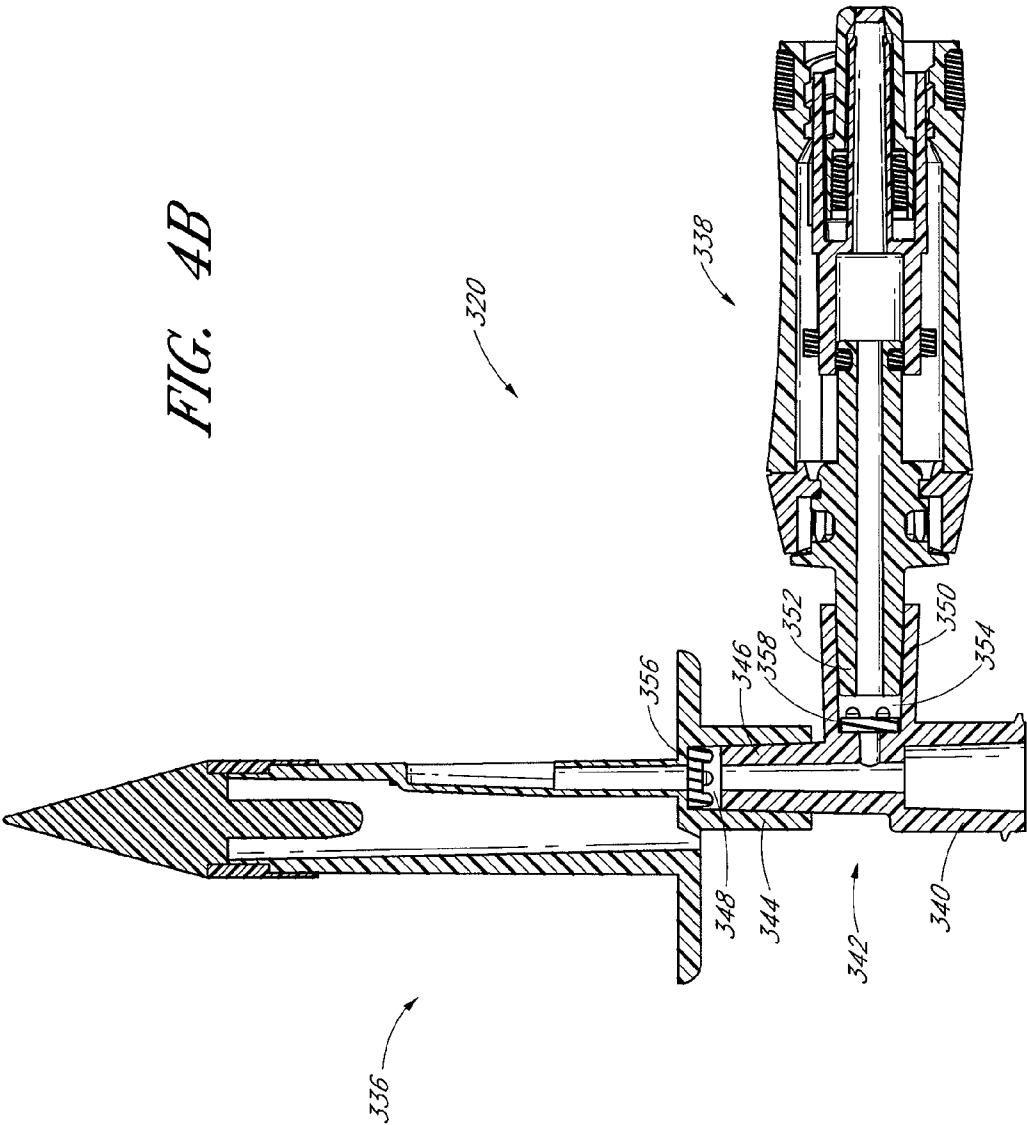


FIG. 3B





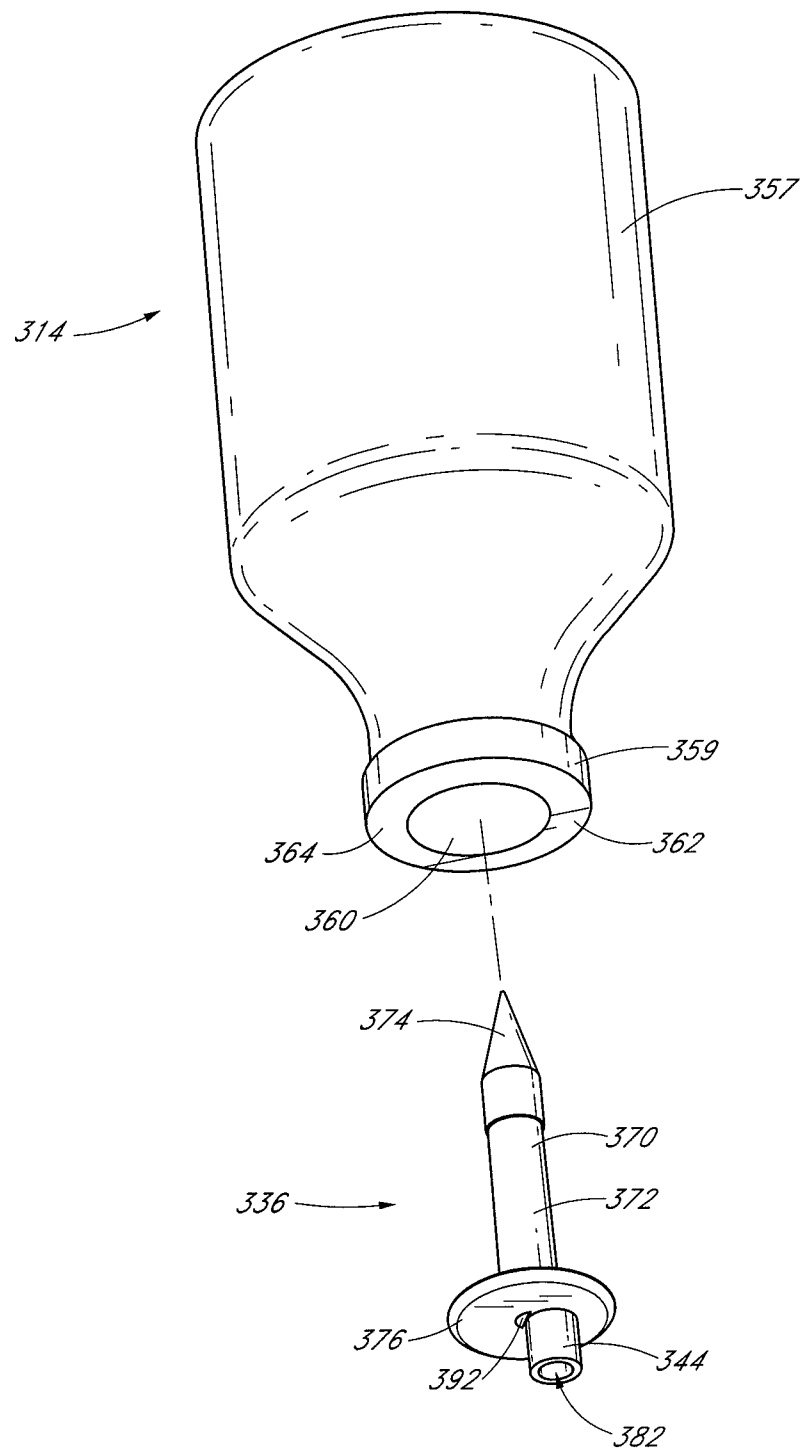


FIG. 5A

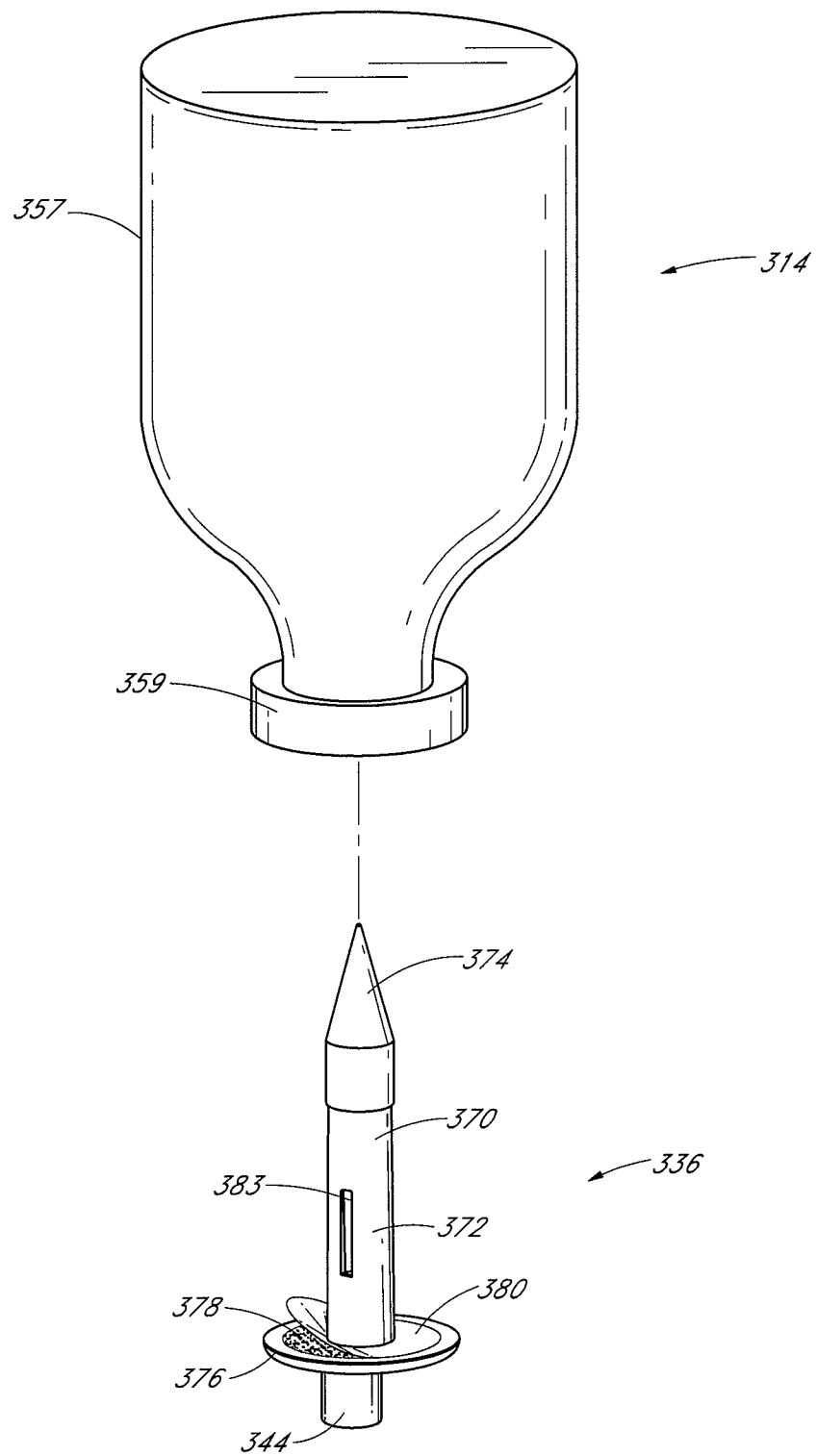
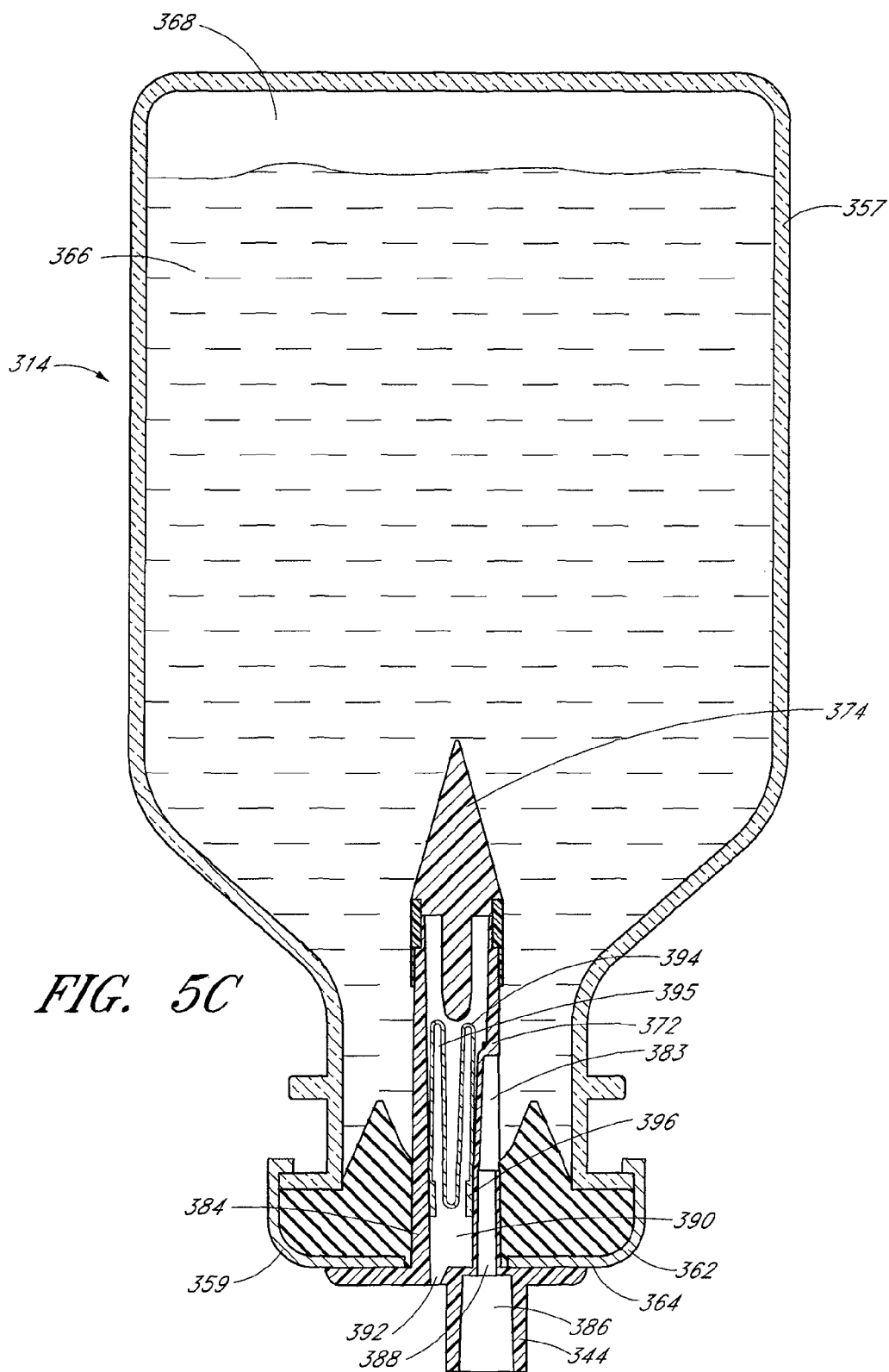
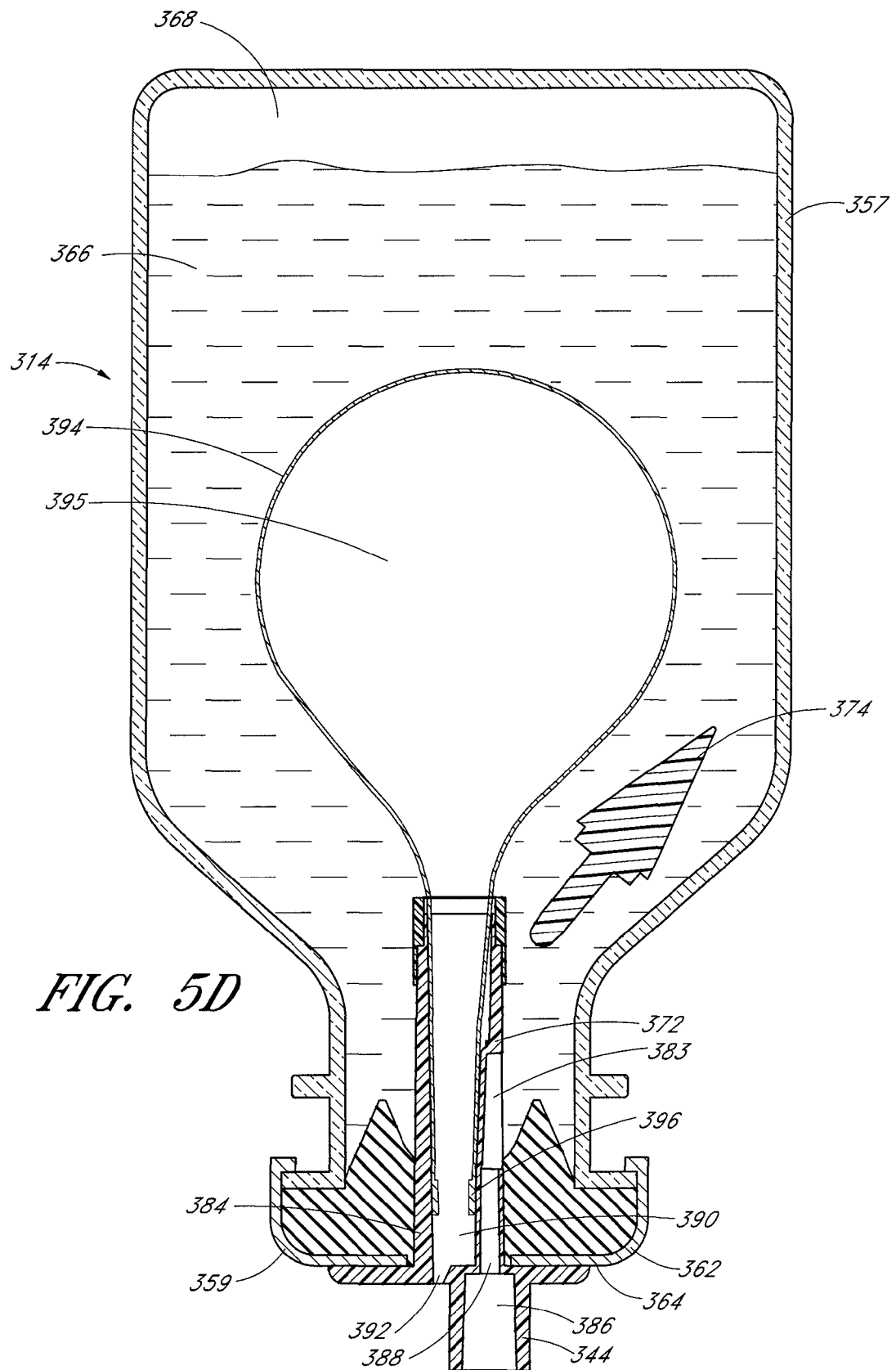


FIG. 5B





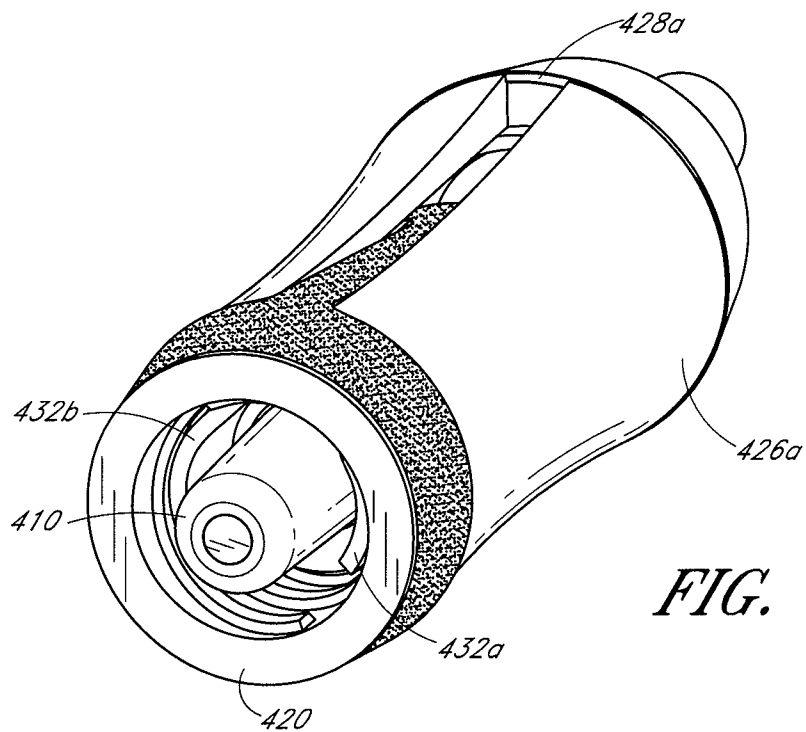


FIG. 6A

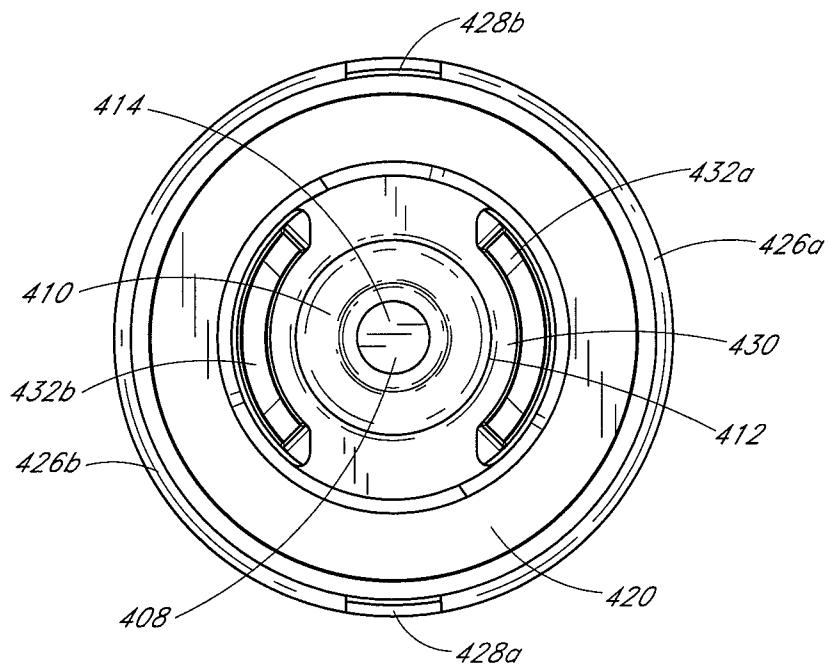


FIG. 6C

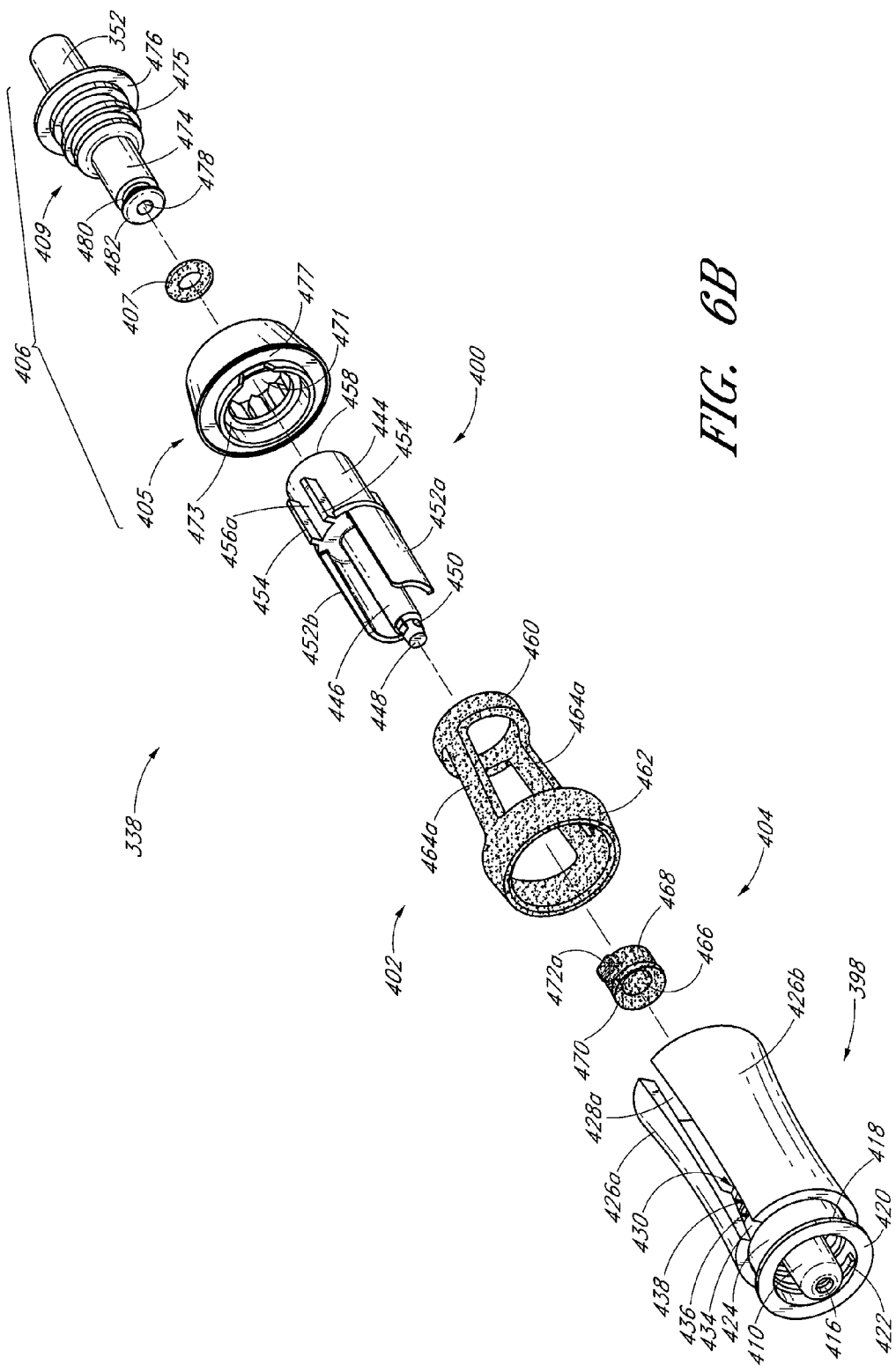


FIG. 6B

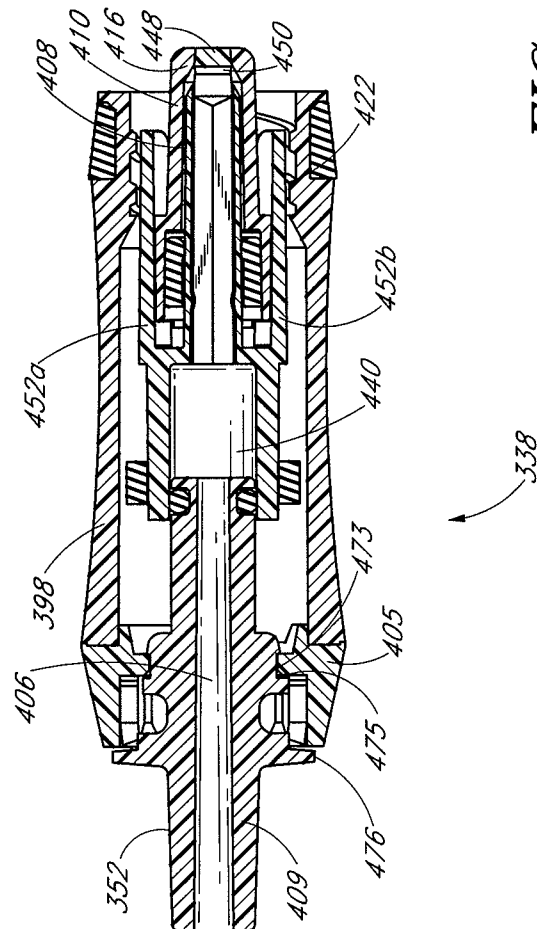
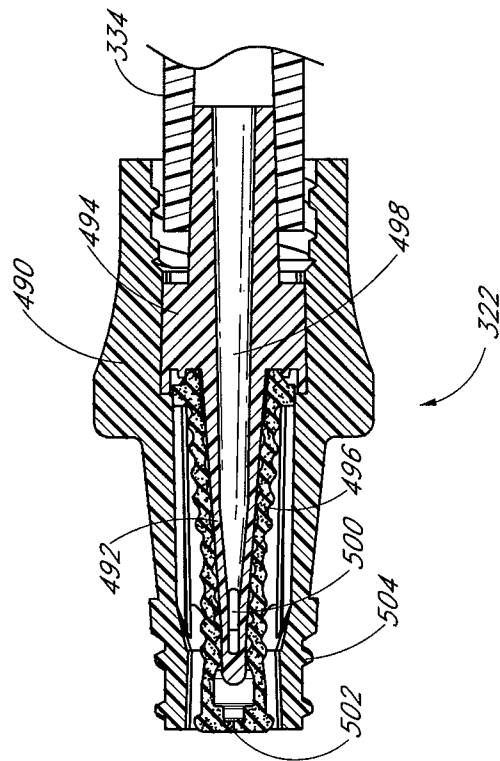


FIG. 6D

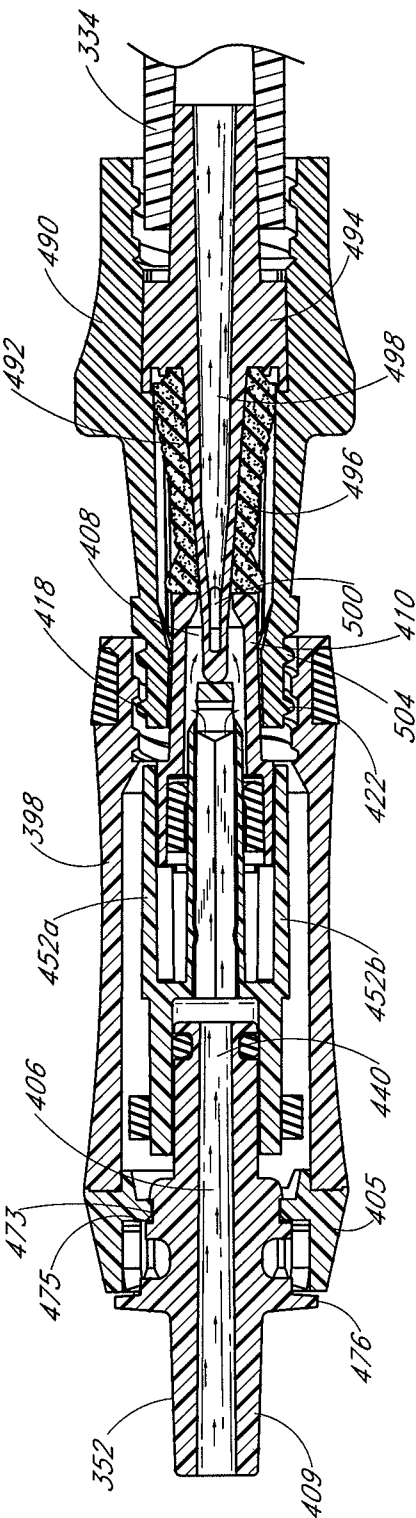
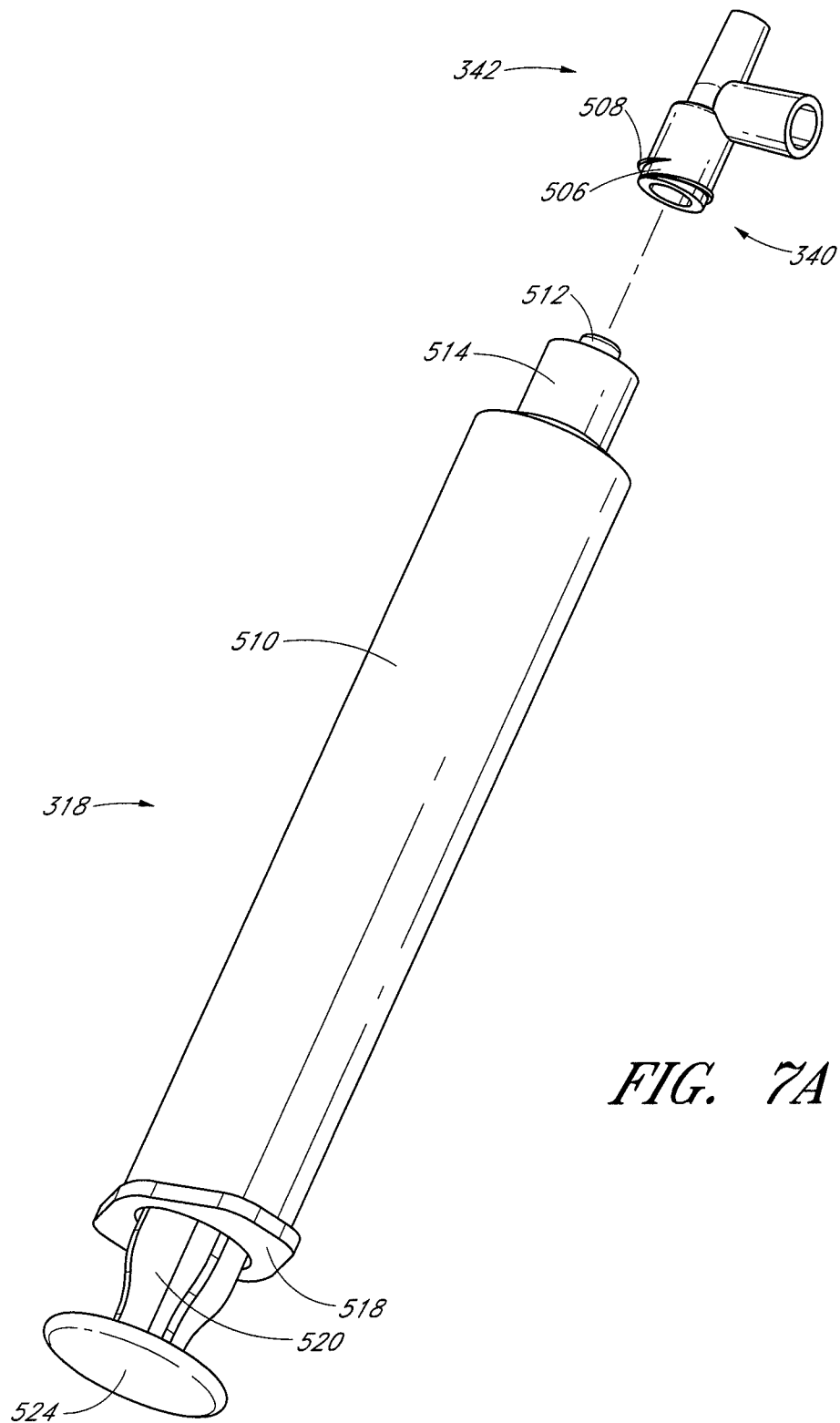
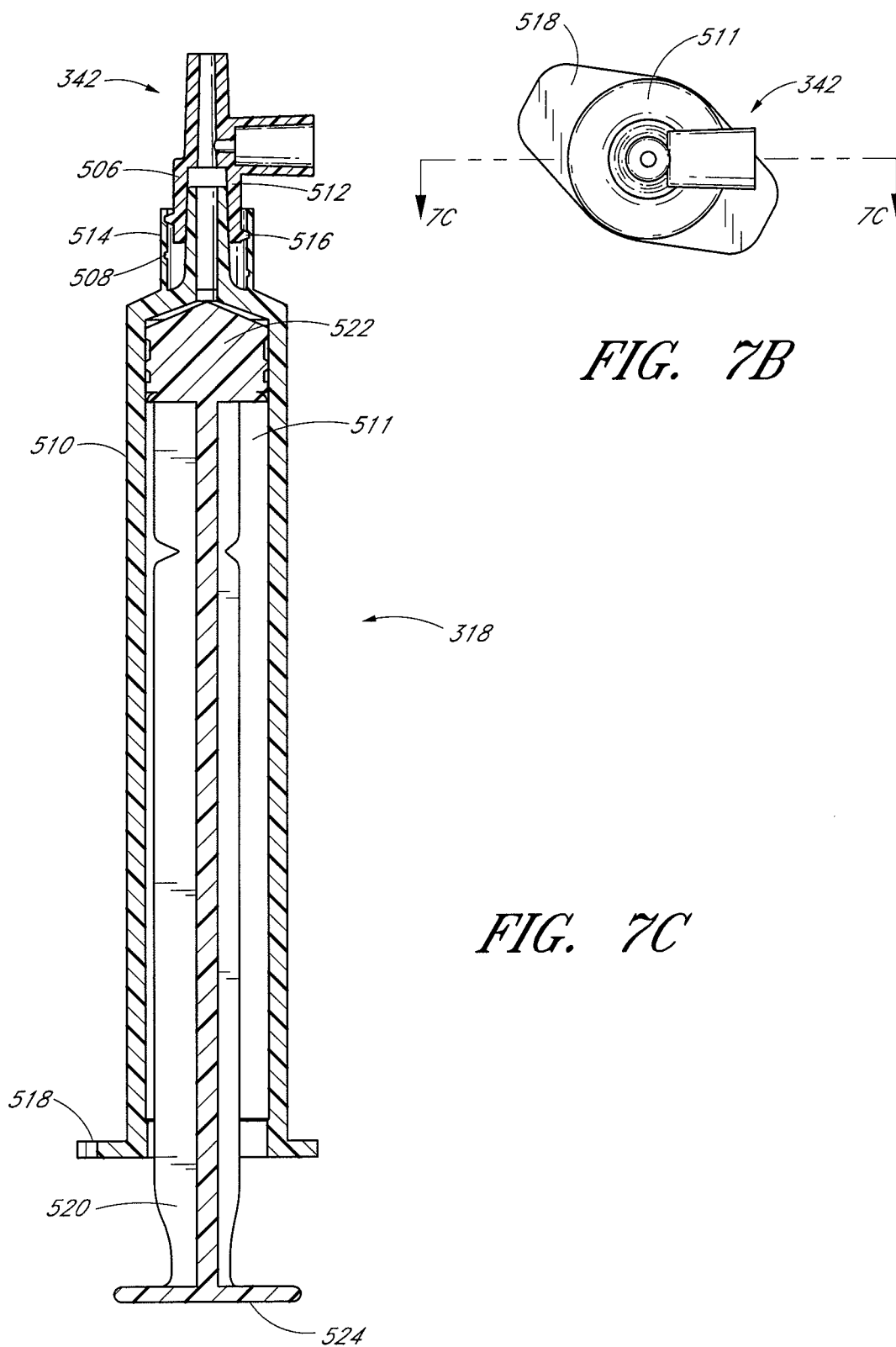


FIG. 6E





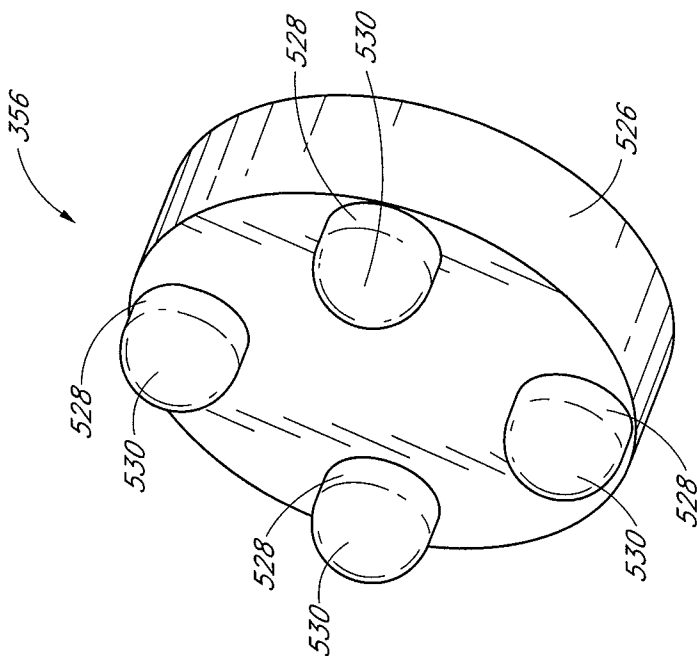


FIG. 8A

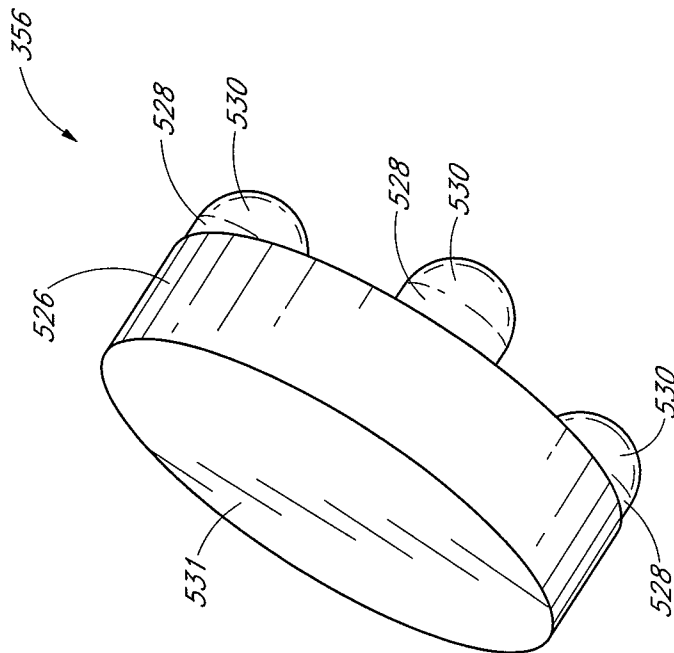


FIG. 8B

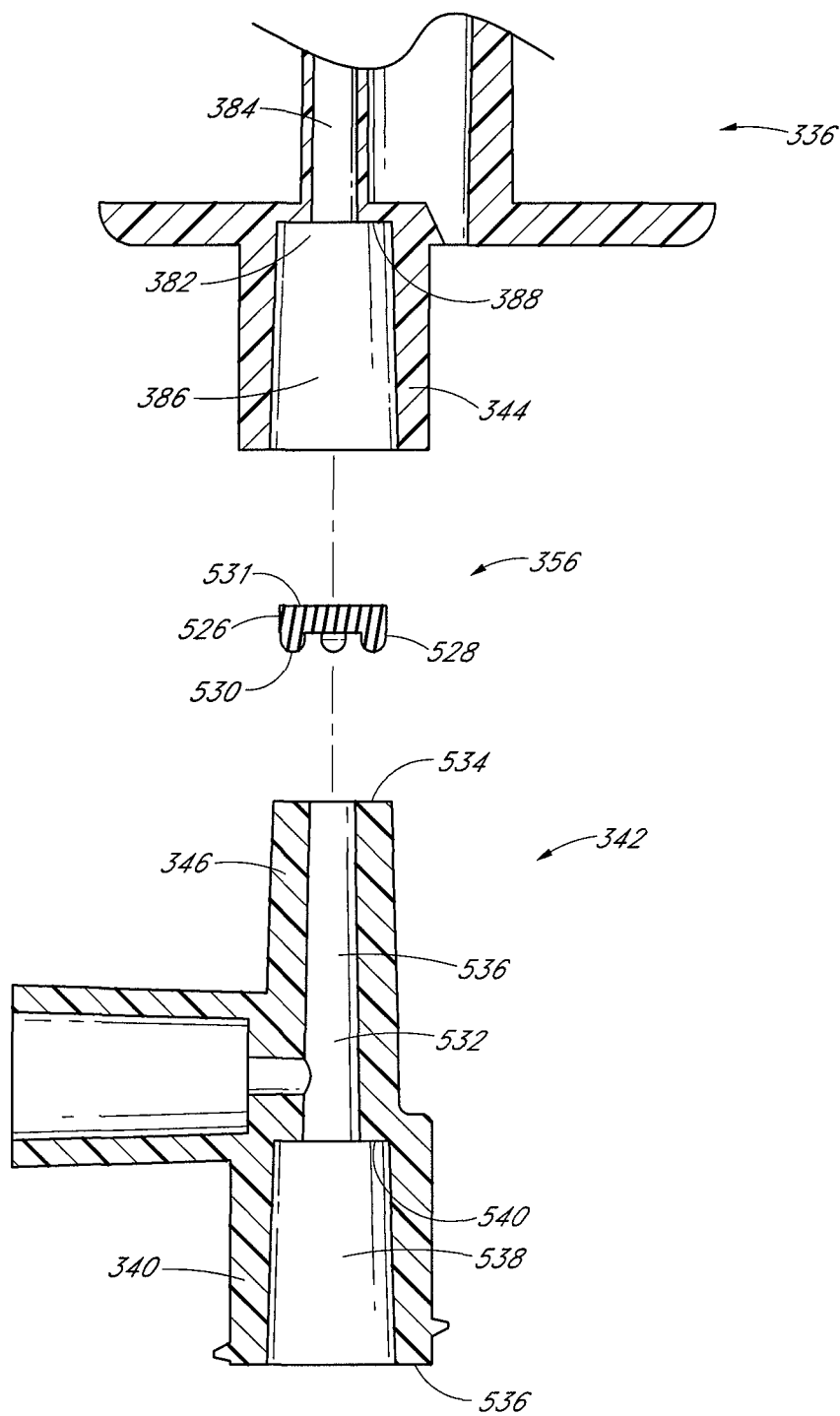
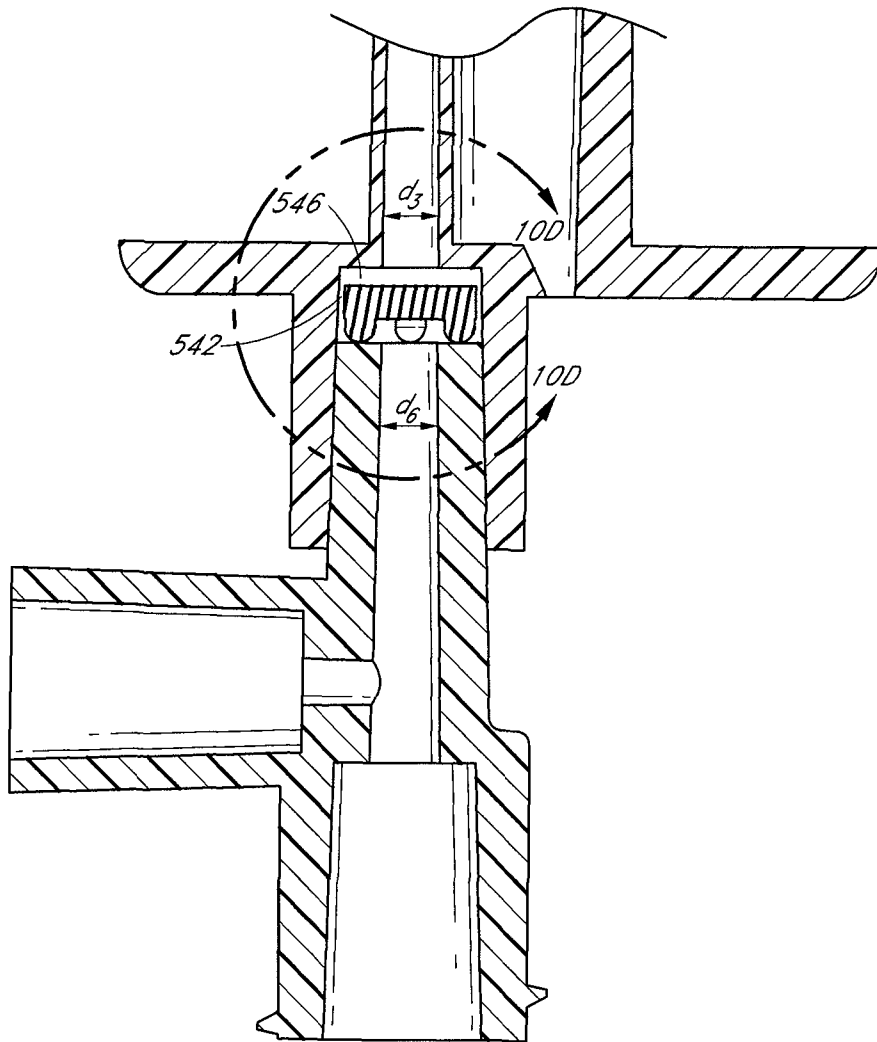


FIG. 9A

*FIG. 9B*

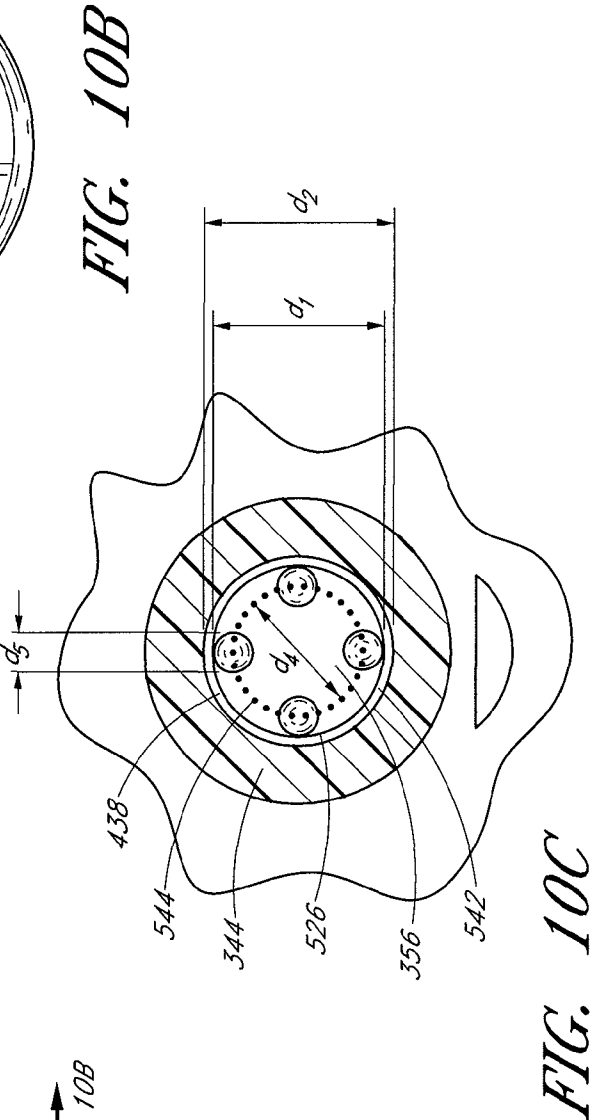
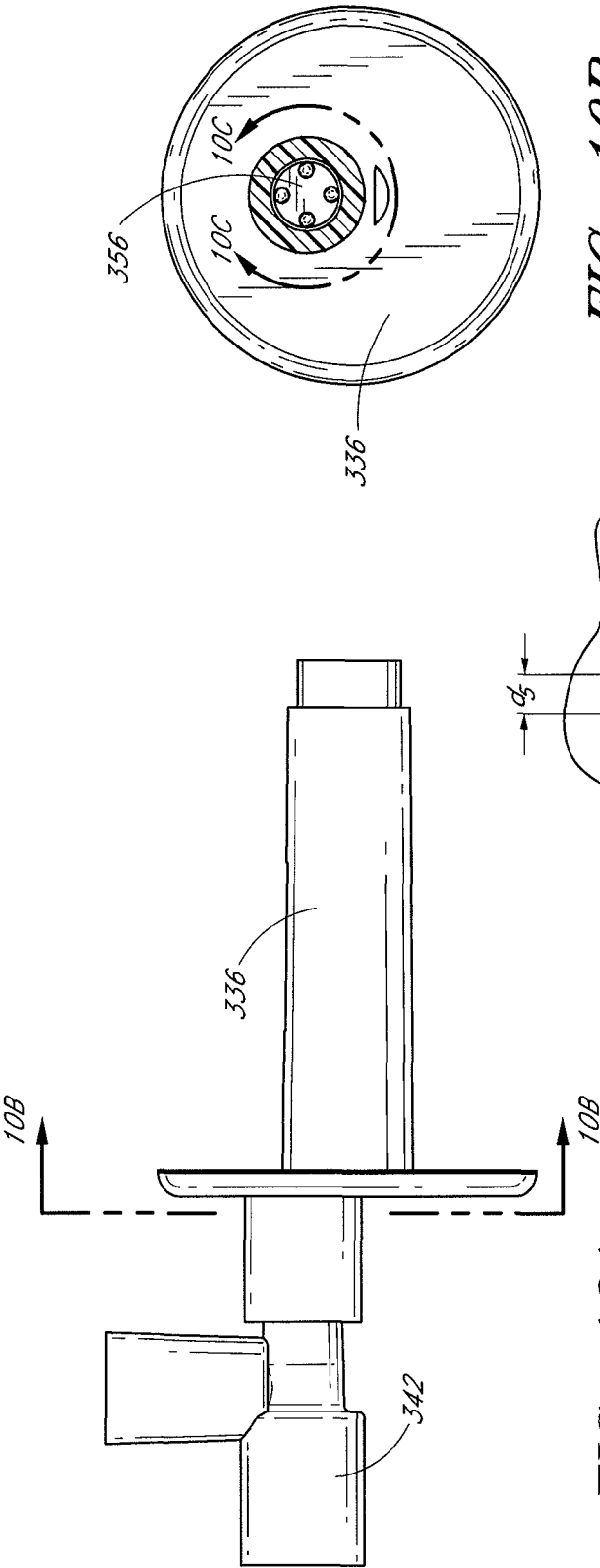
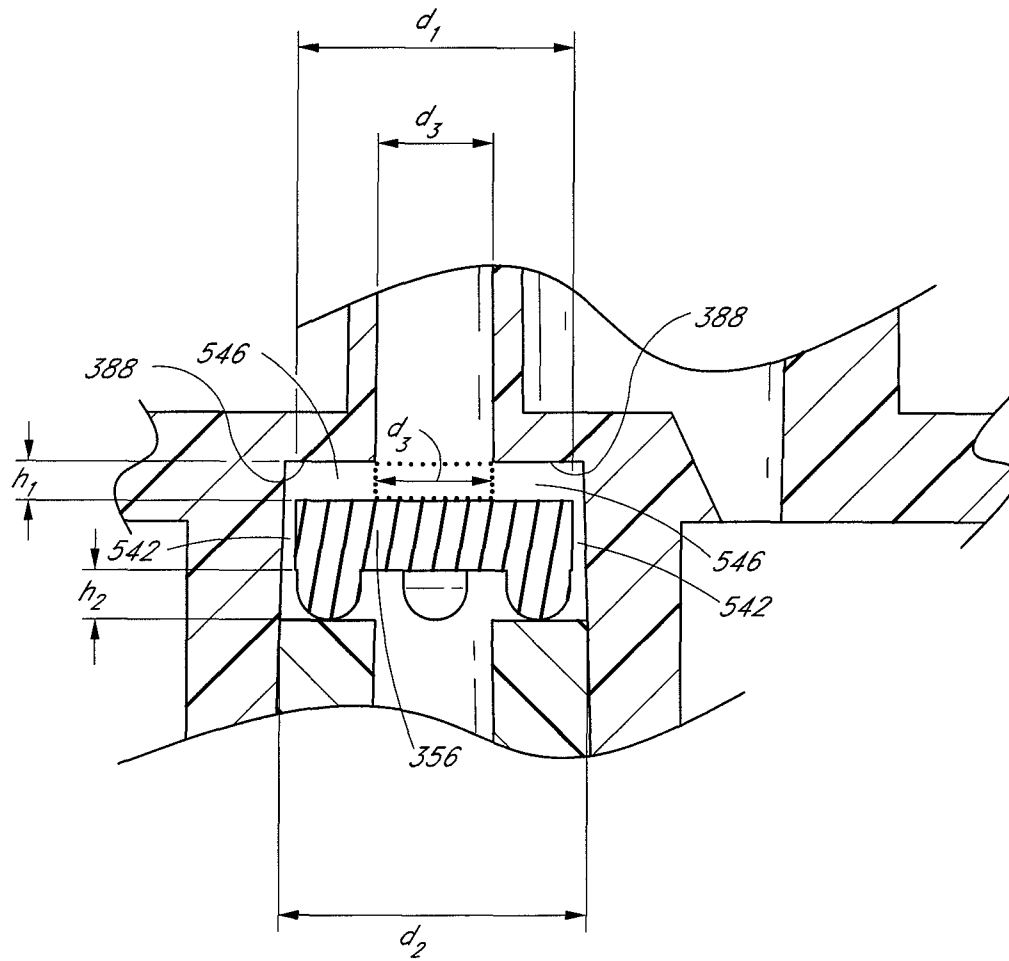
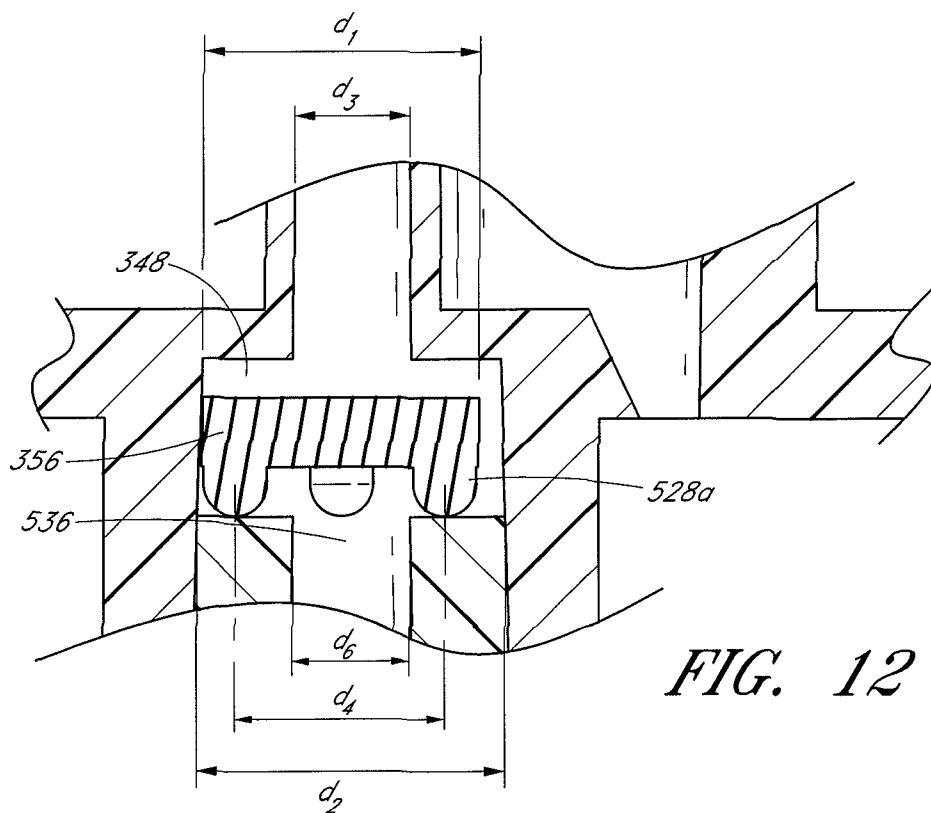
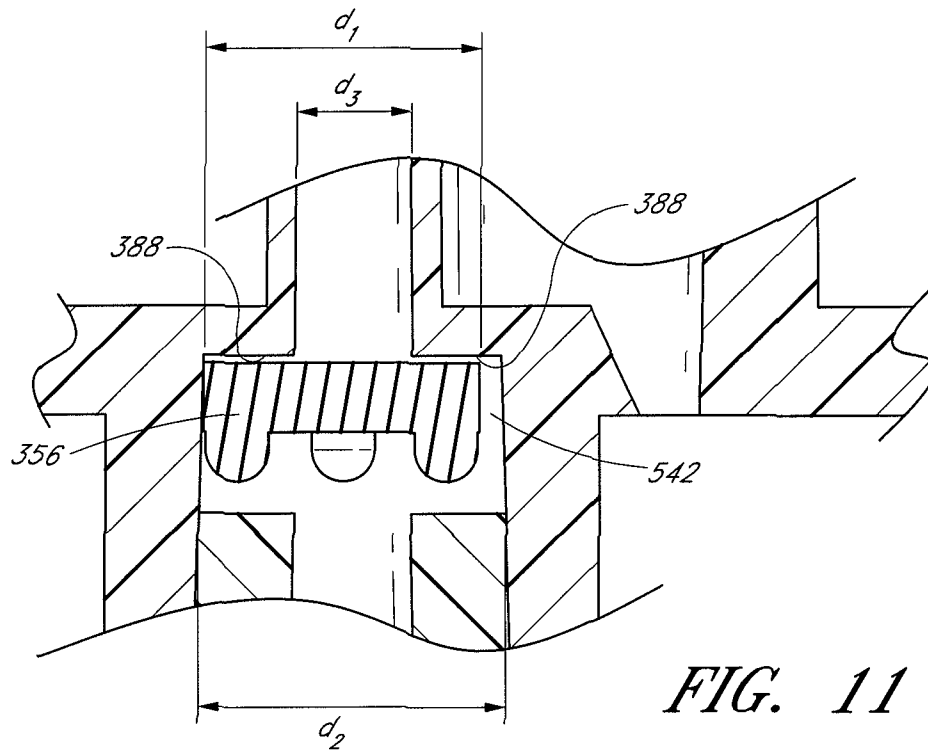


FIG. 10C

*FIG. 10D*



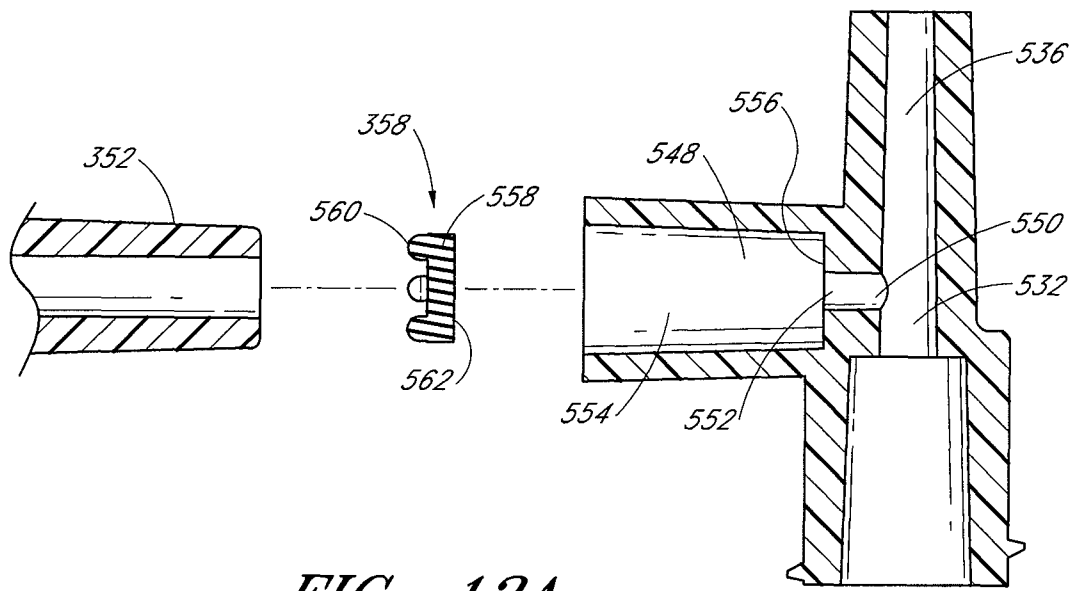


FIG. 13A

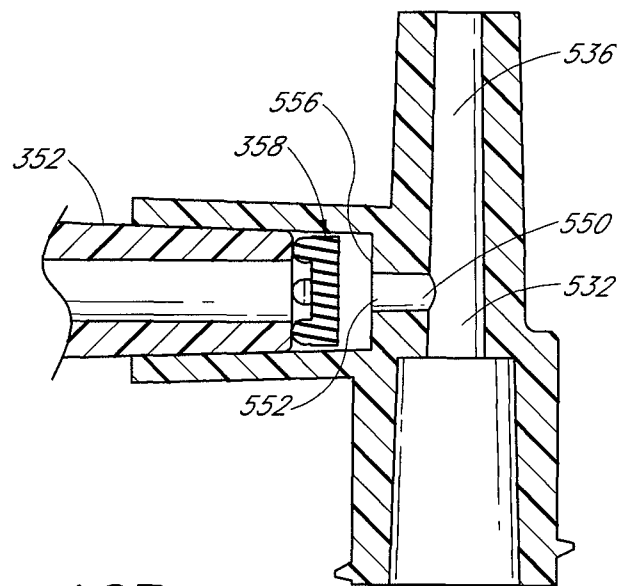


FIG. 13B

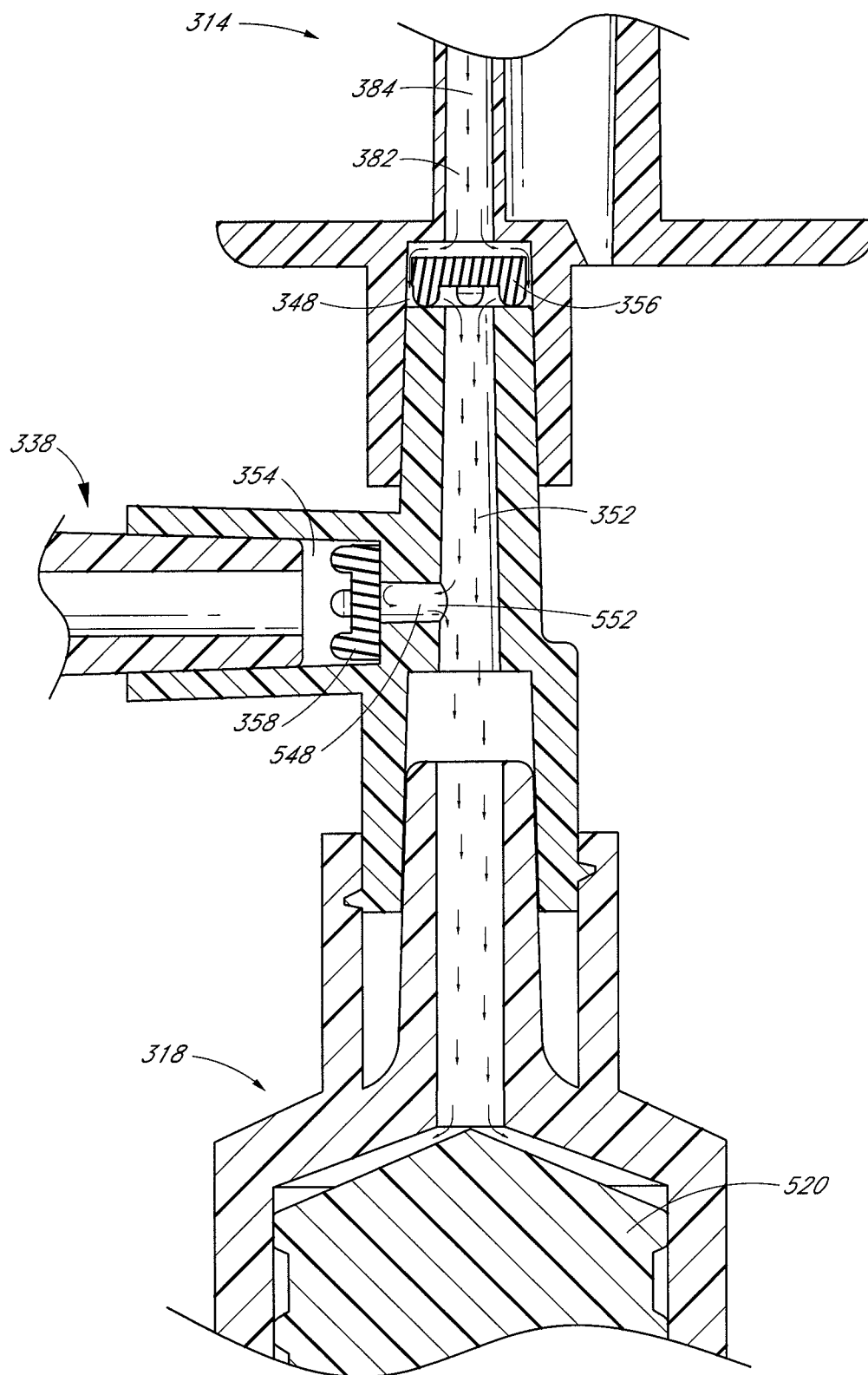
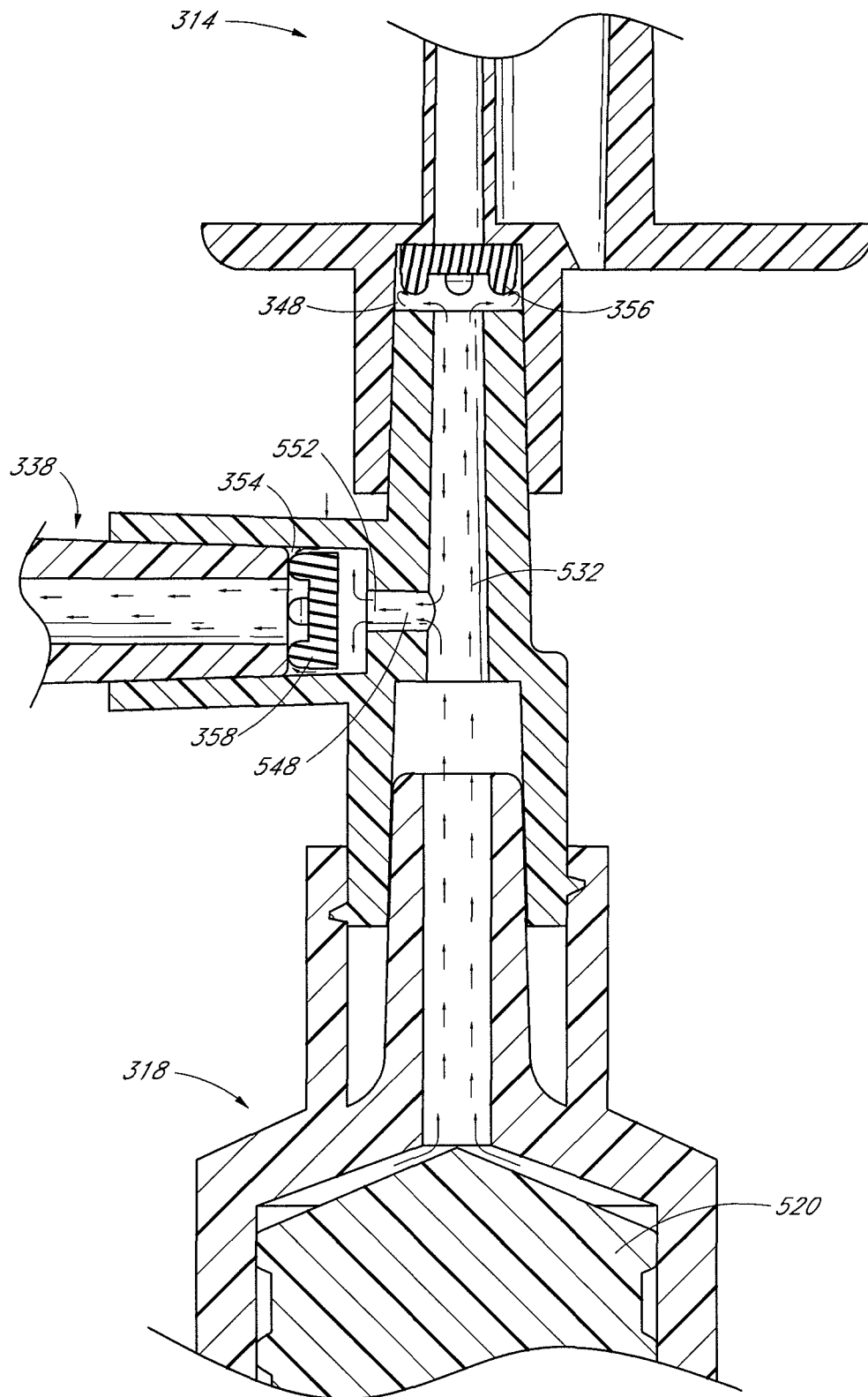
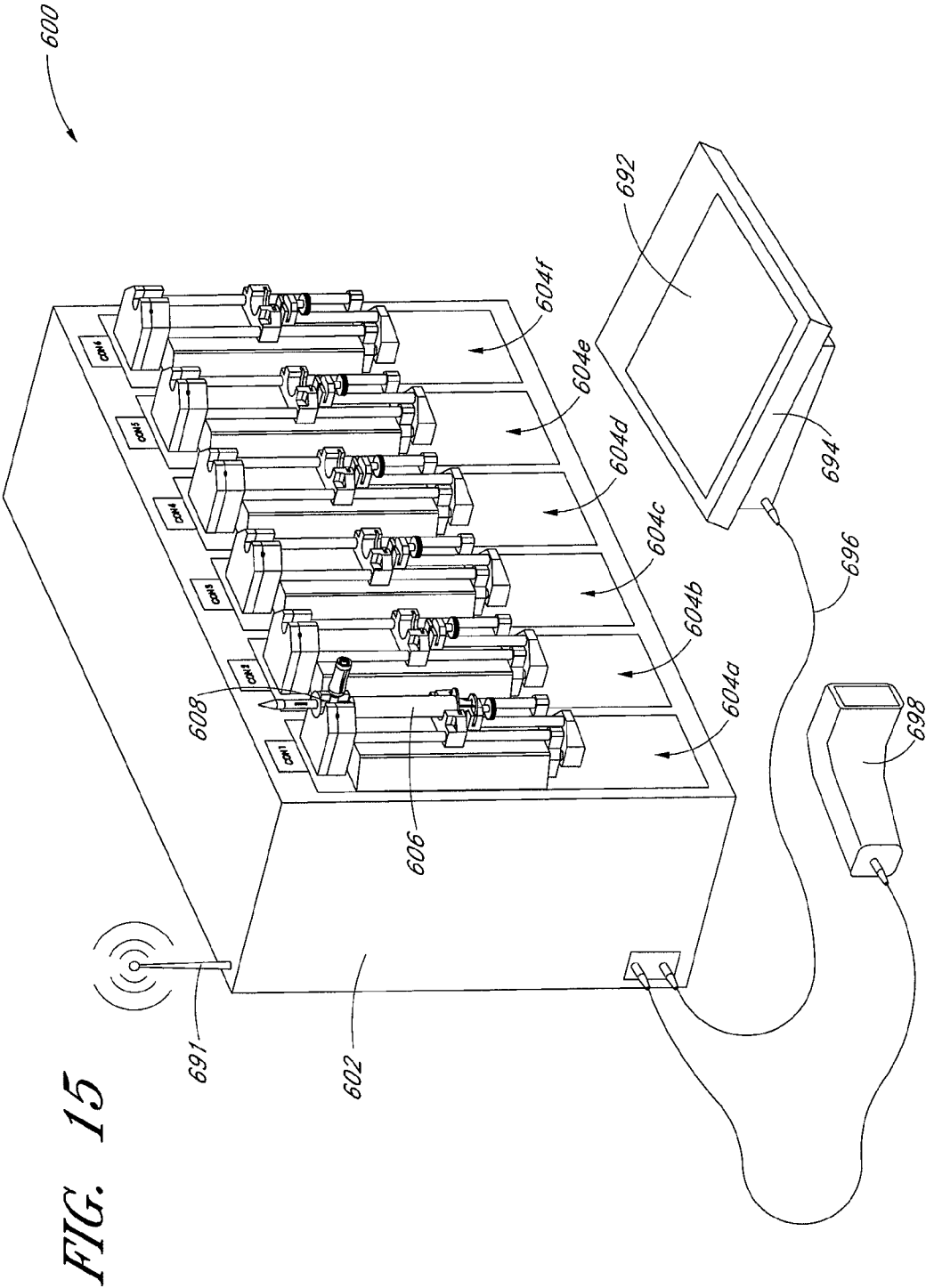


FIG. 14A

*FIG. 14B*



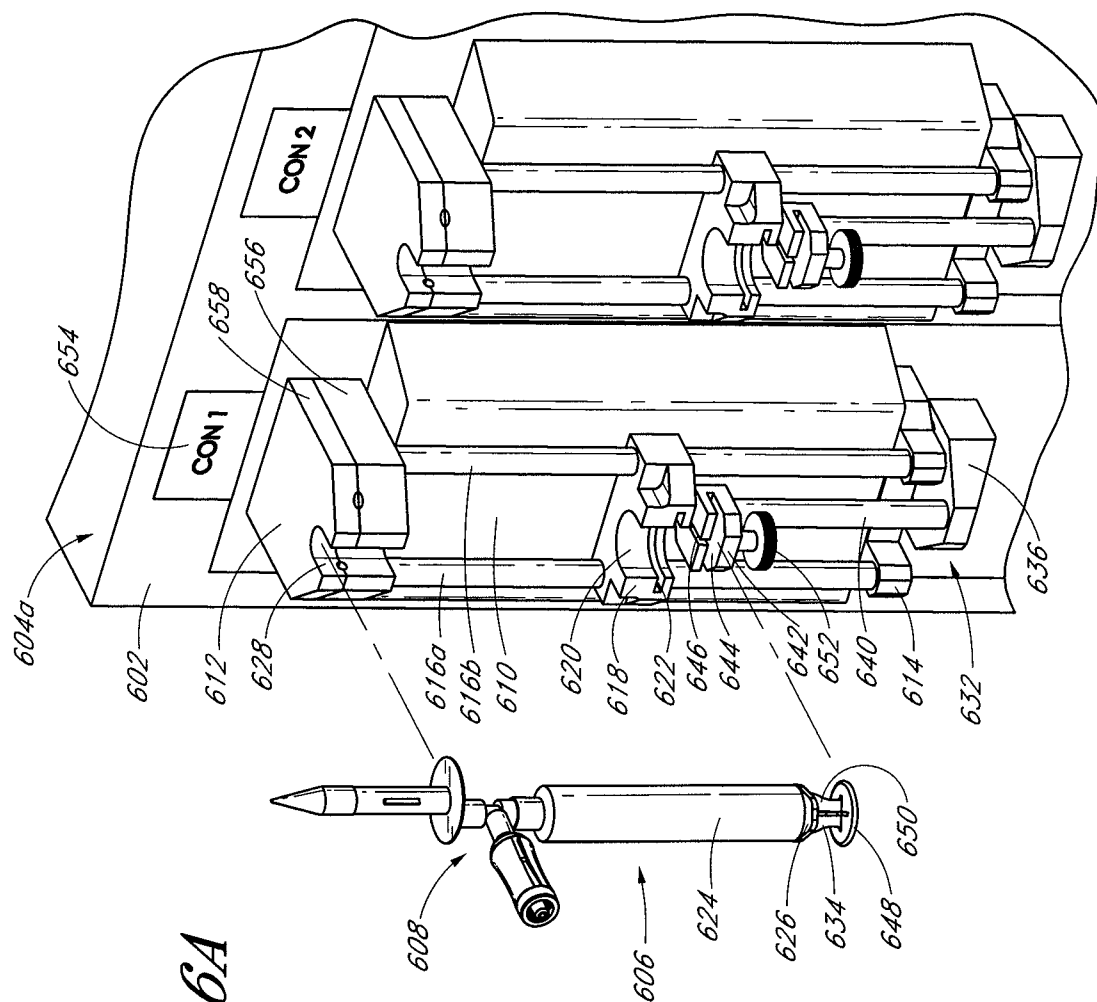
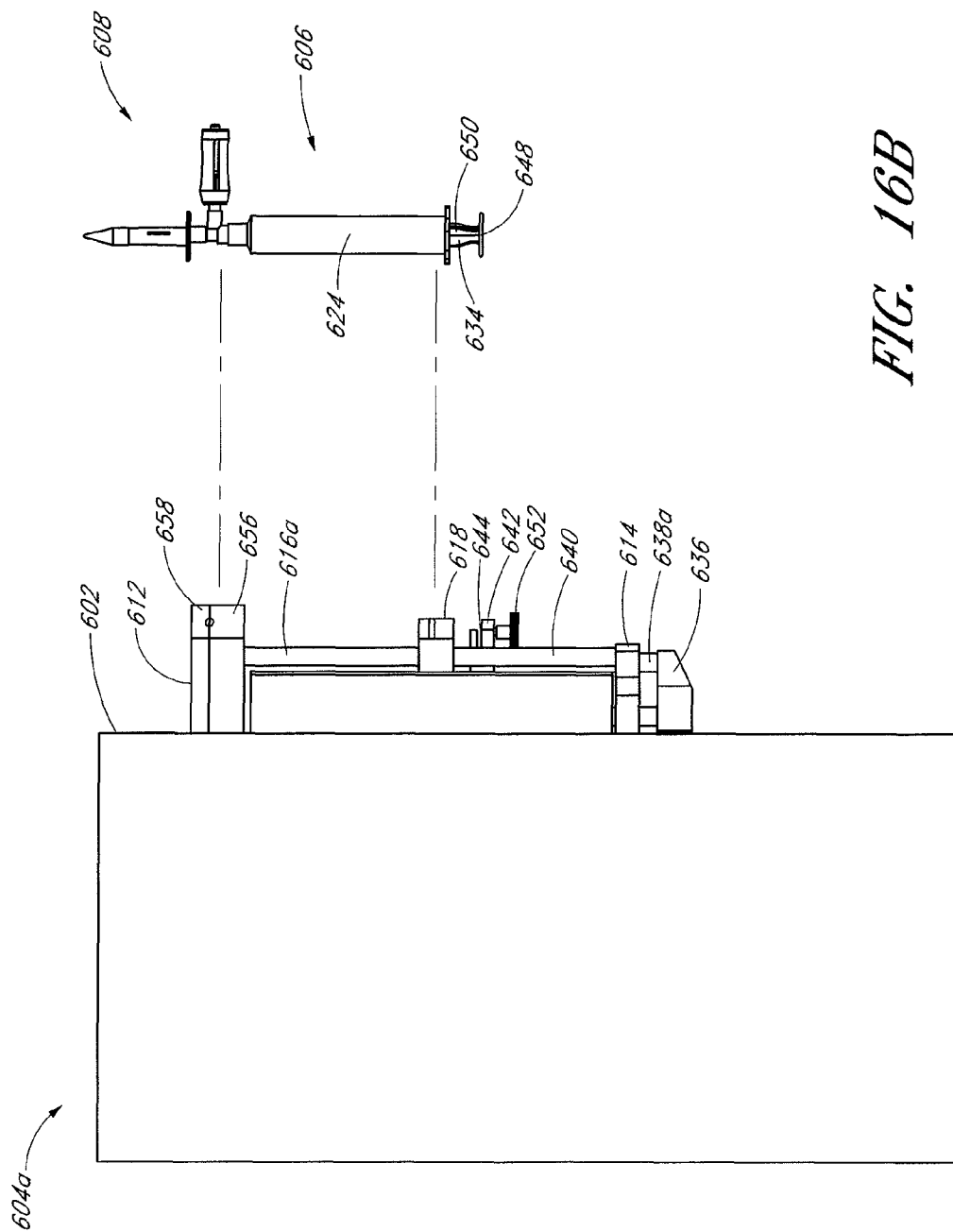


FIG. 16A



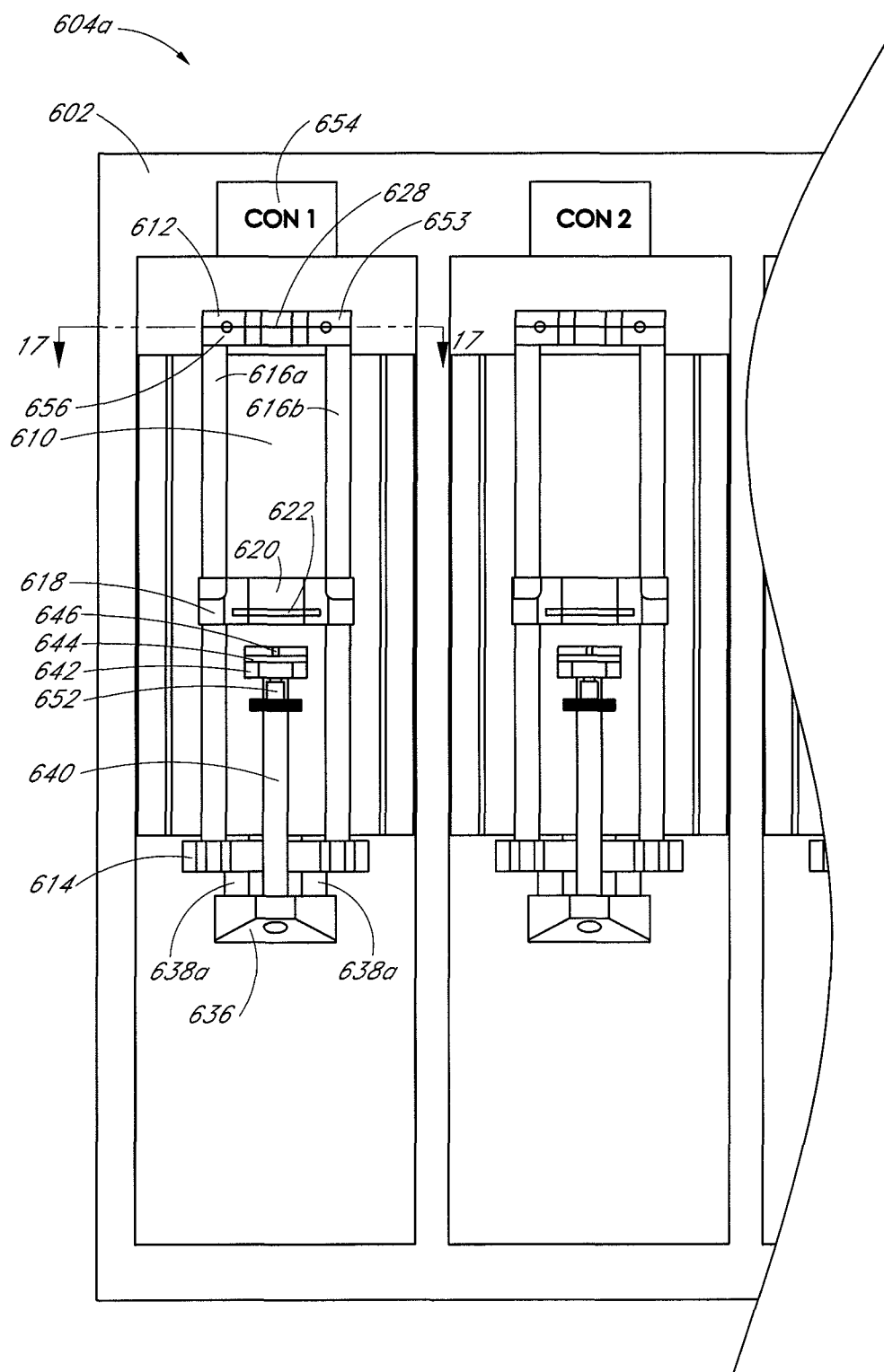


FIG. 16C

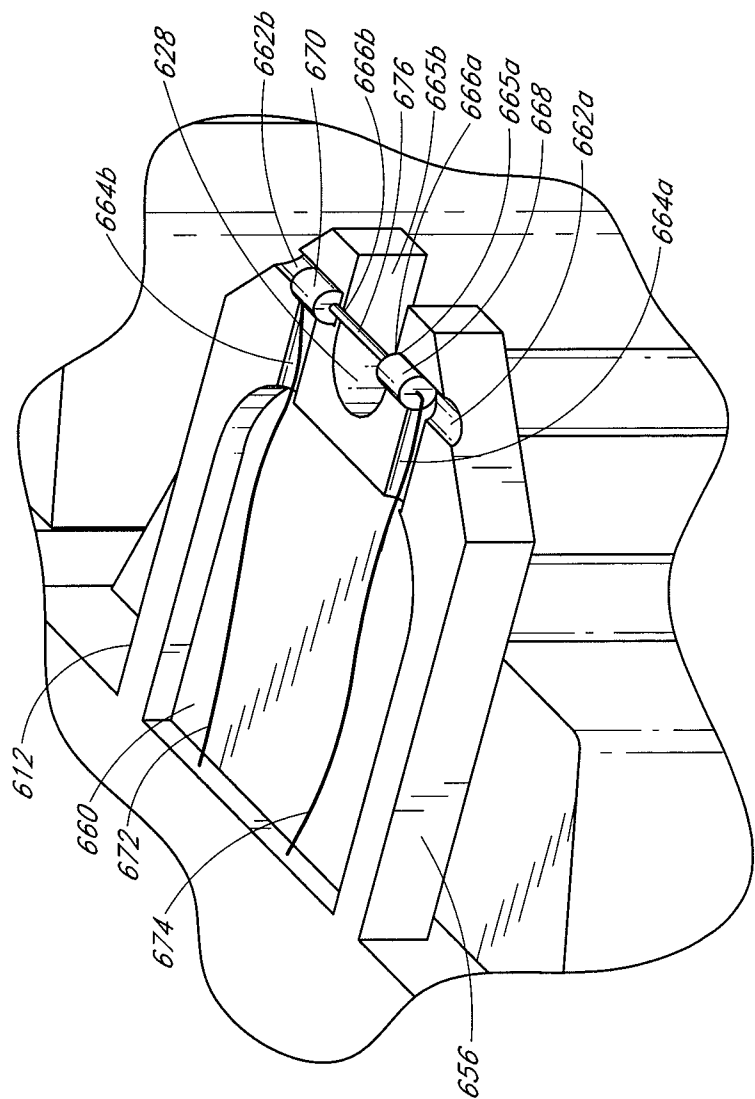


FIG. 17

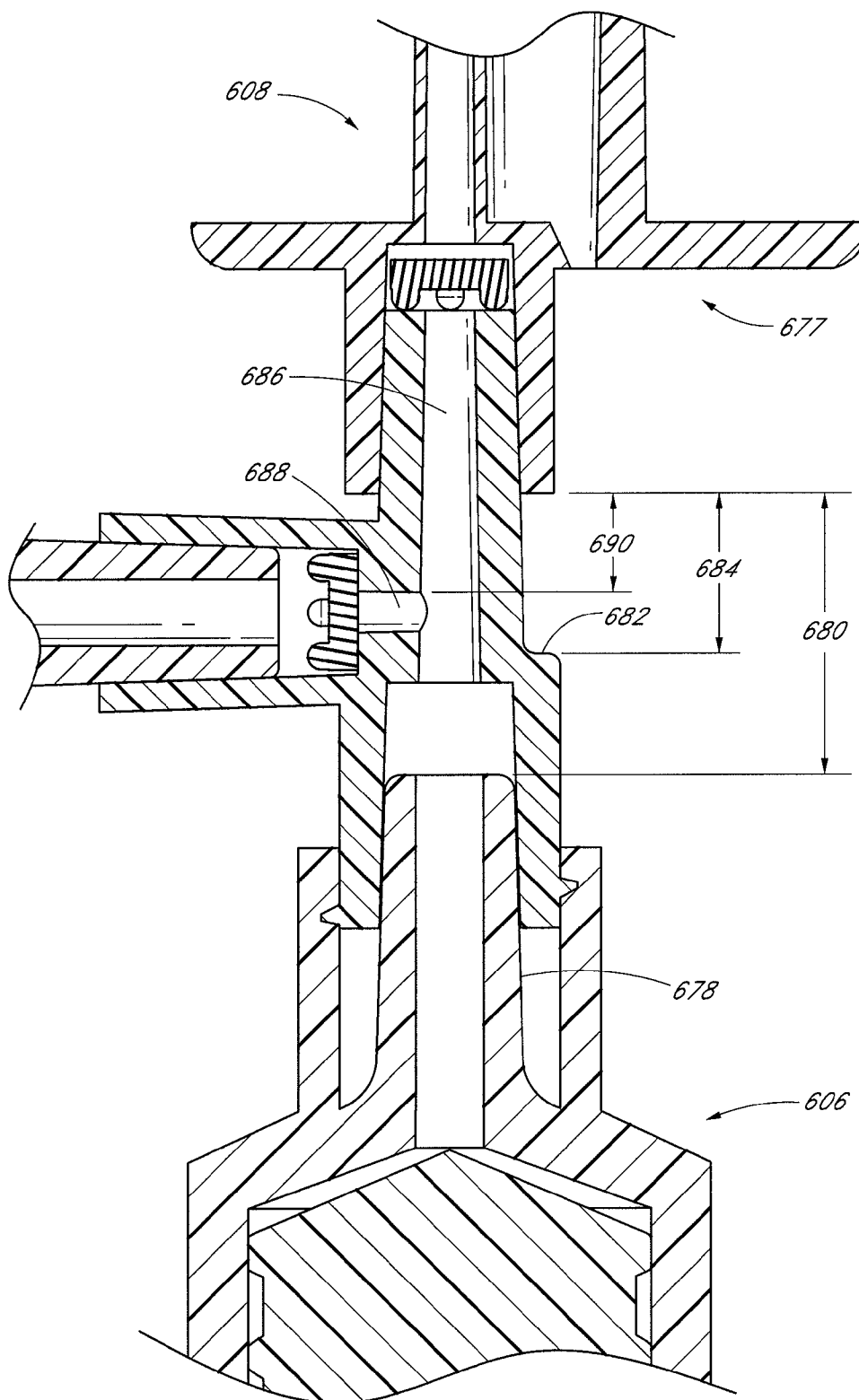


FIG. 18

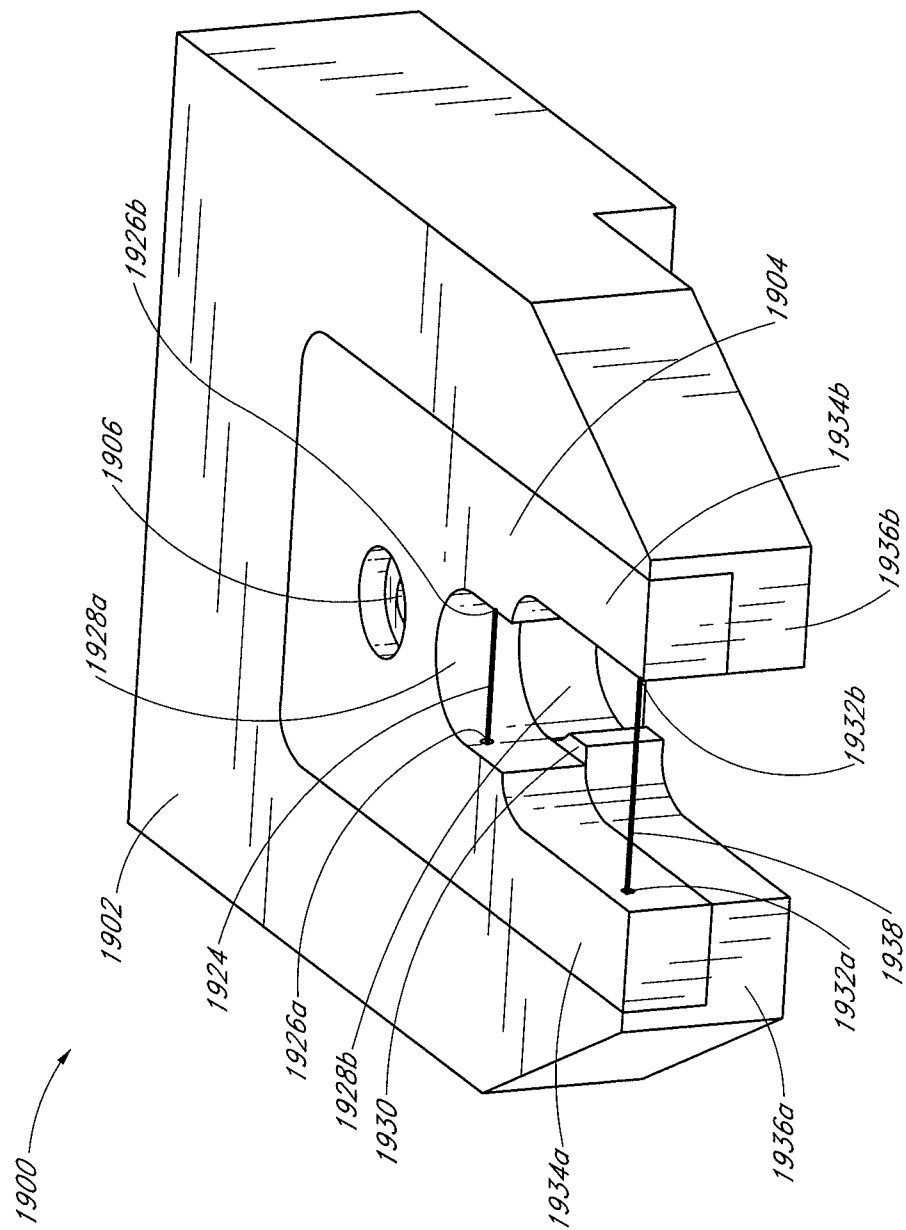


FIG. 19A

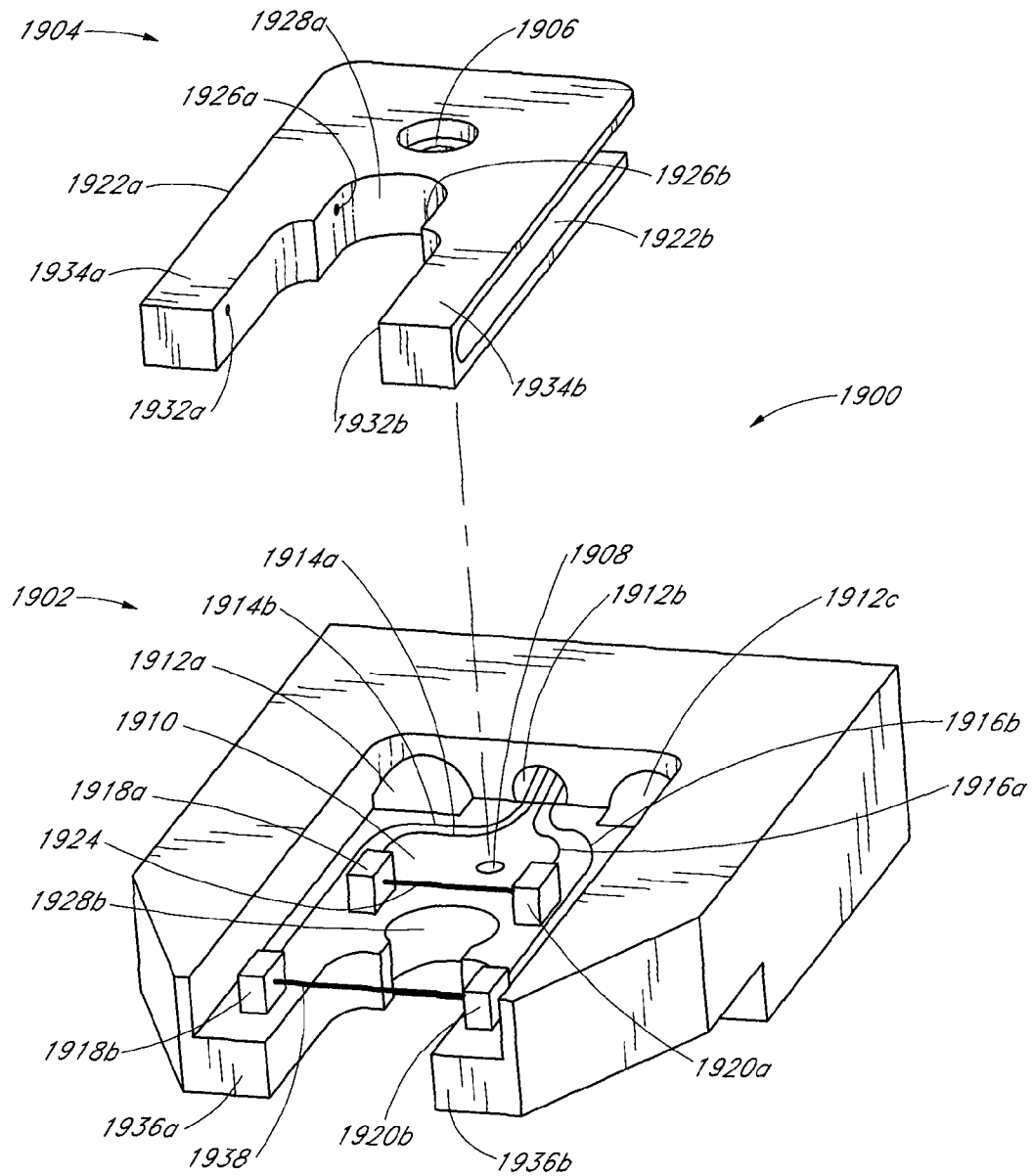


FIG. 19B

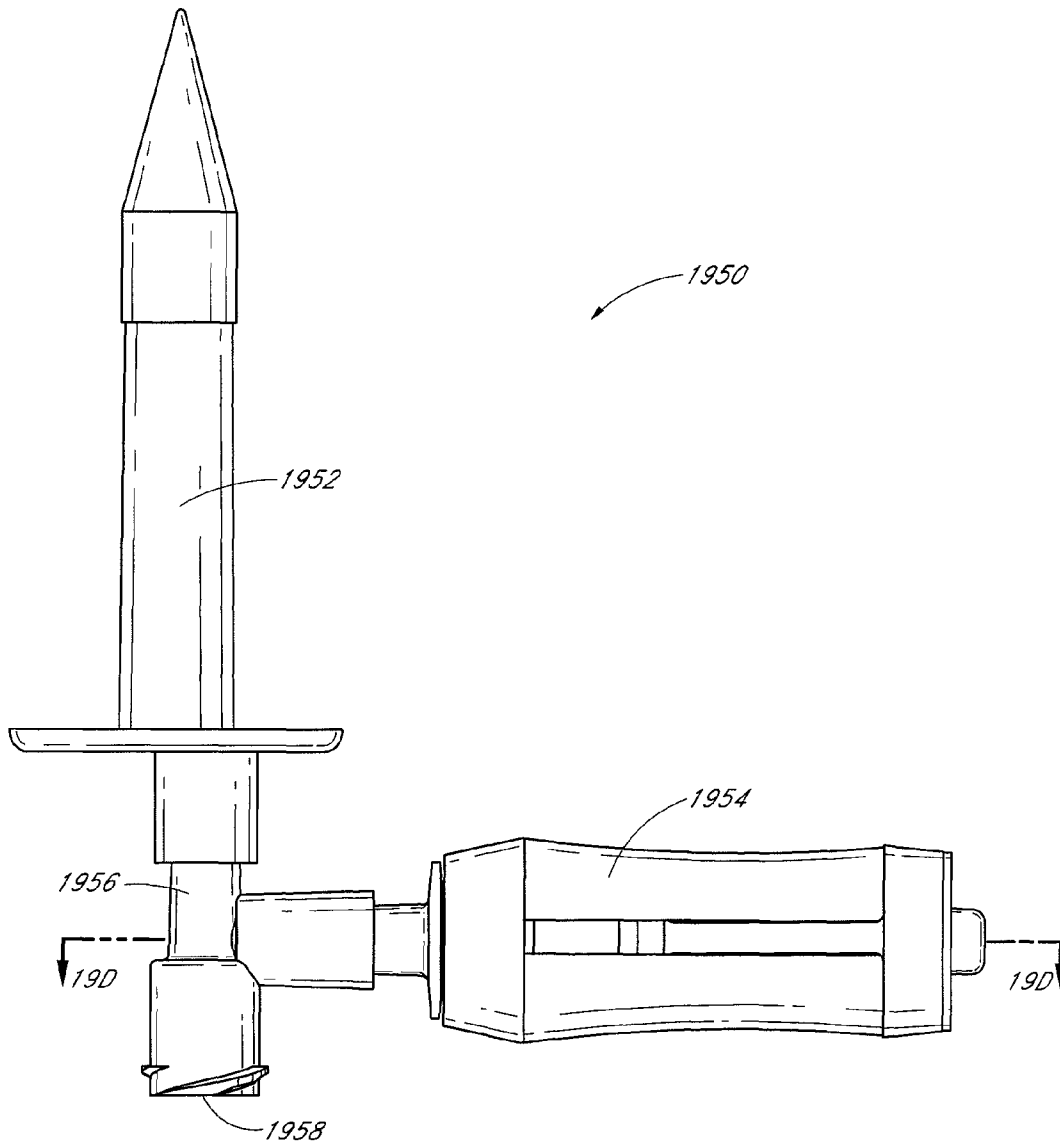


FIG. 19C

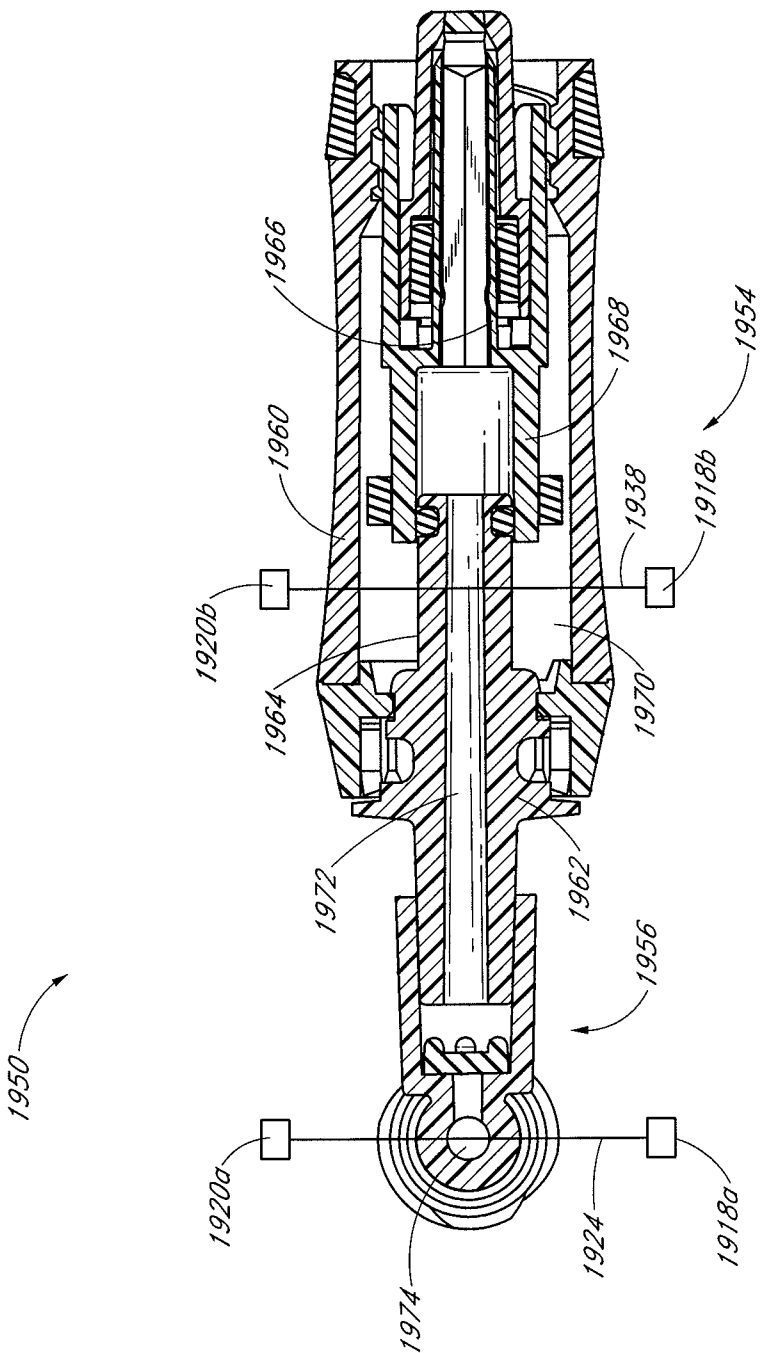


FIG. 19D

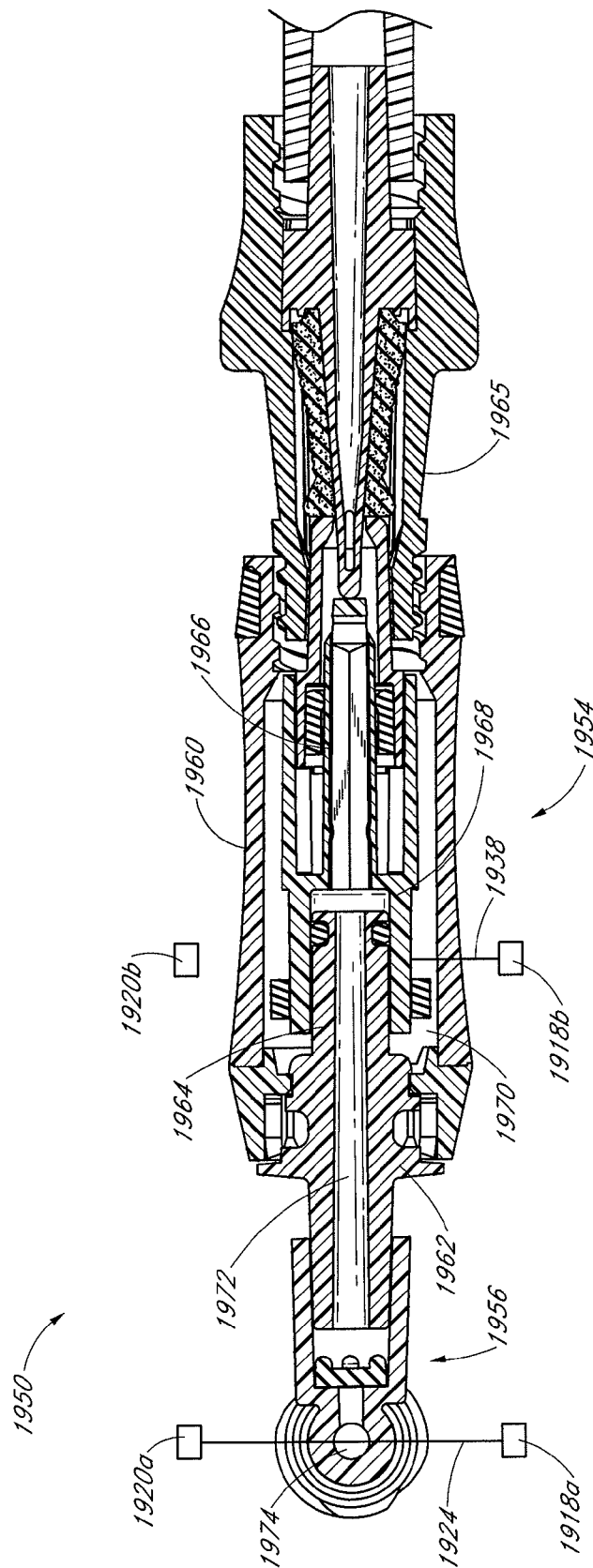
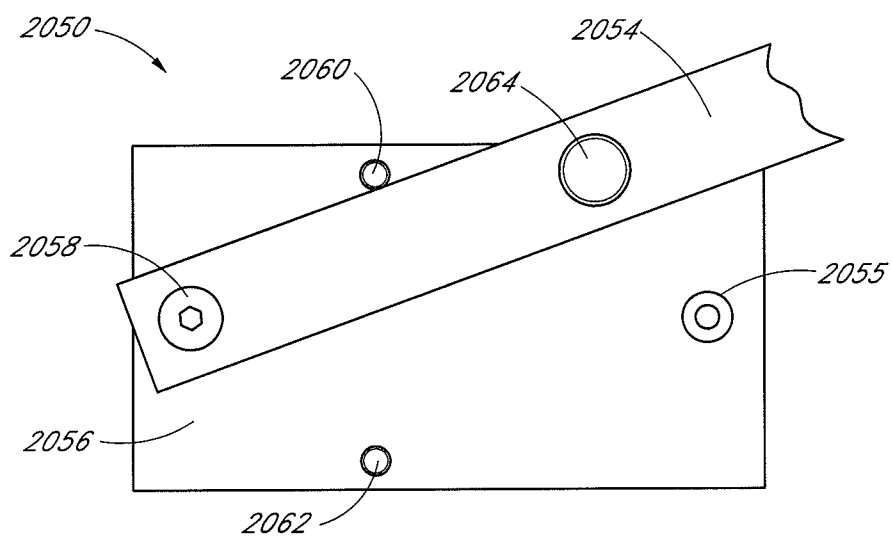
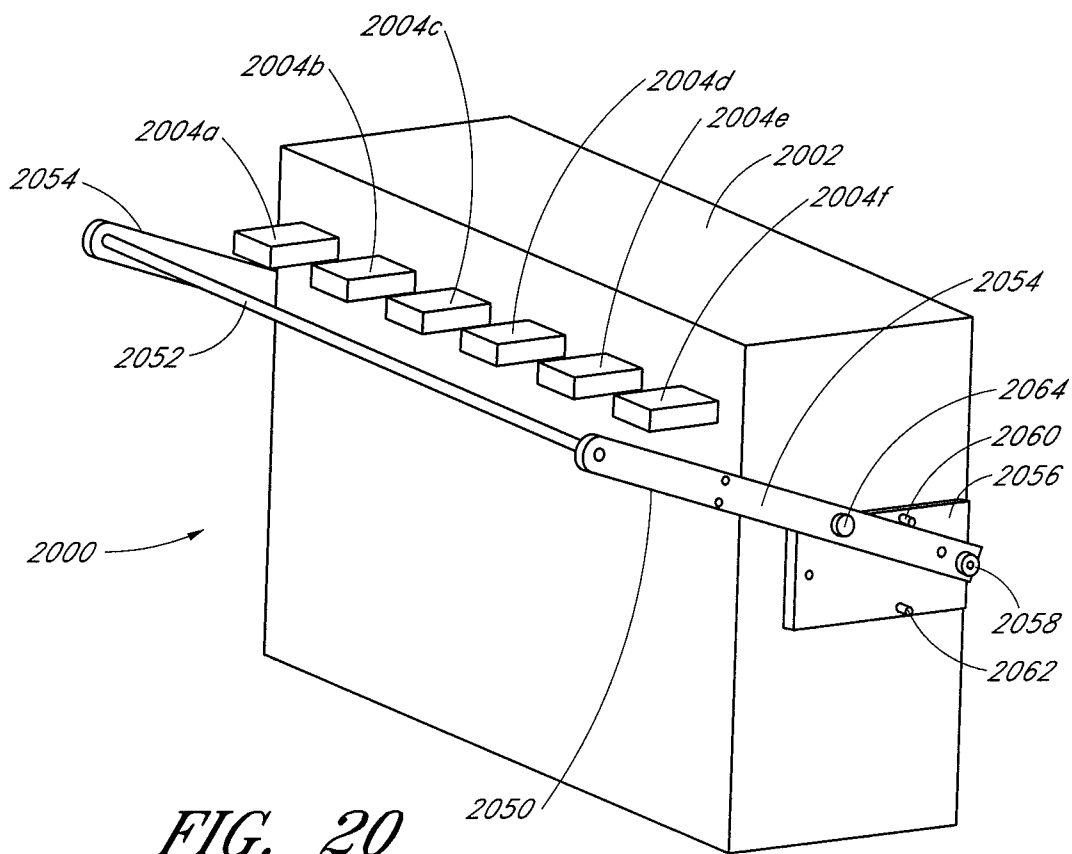


FIG. 19E



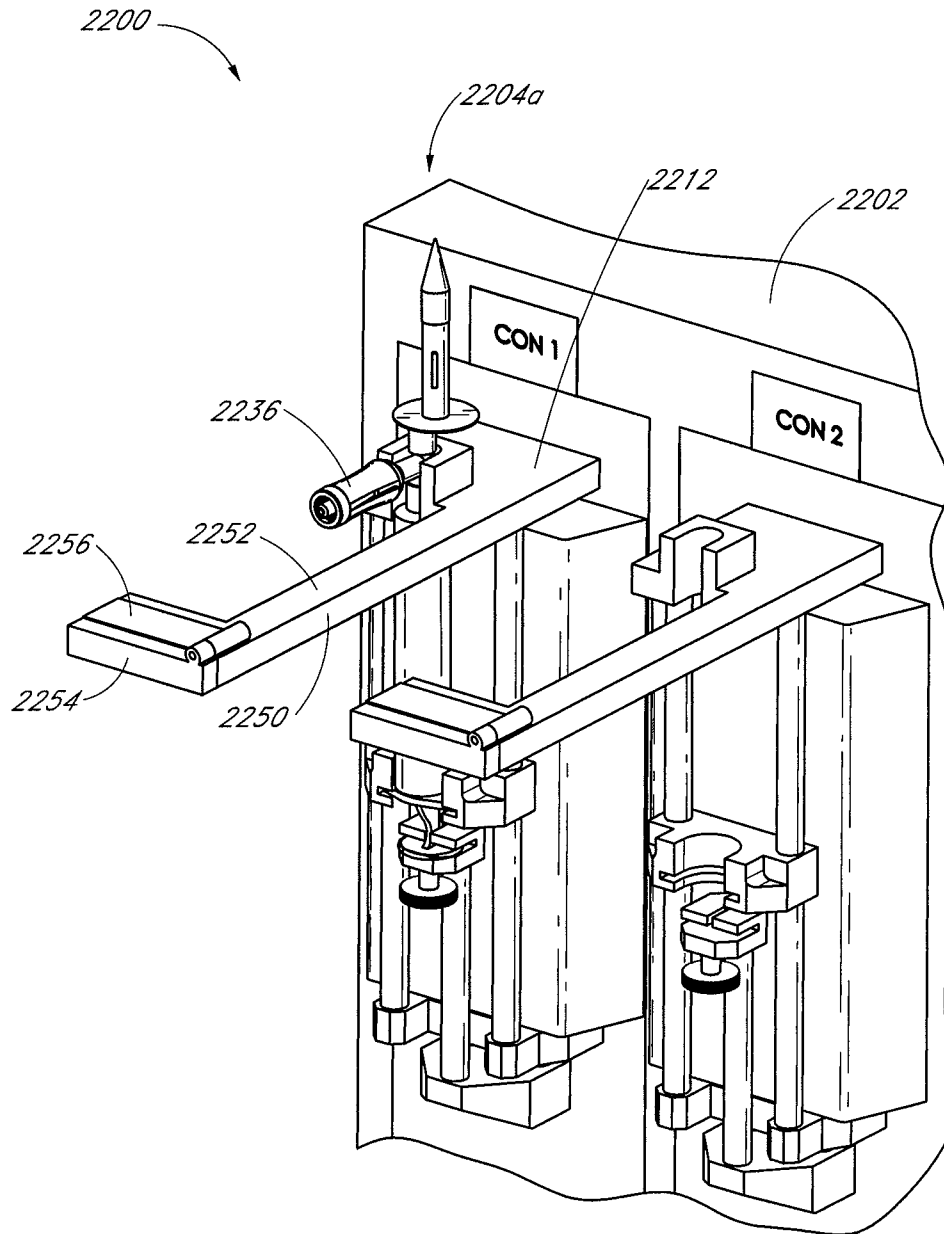
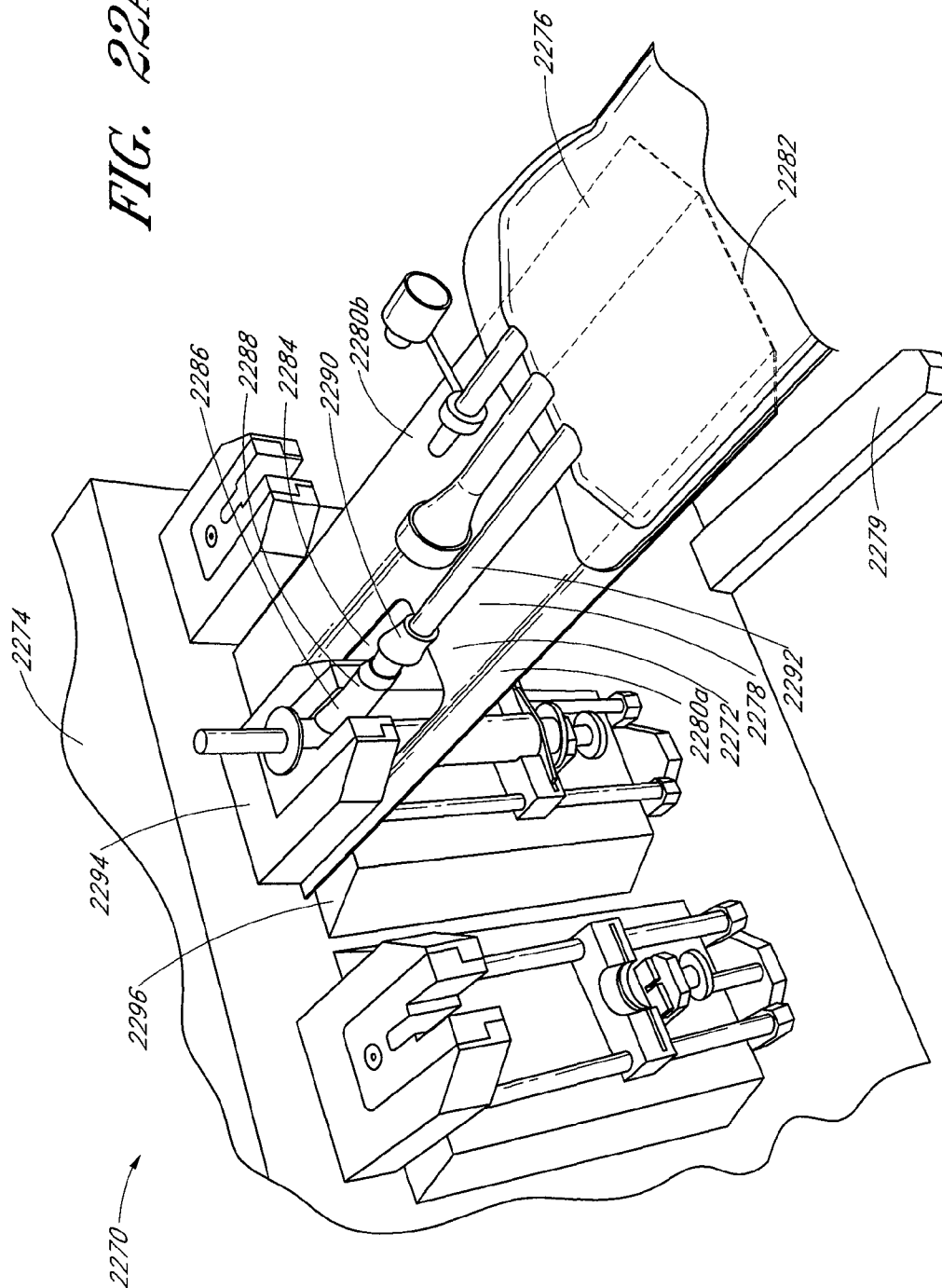
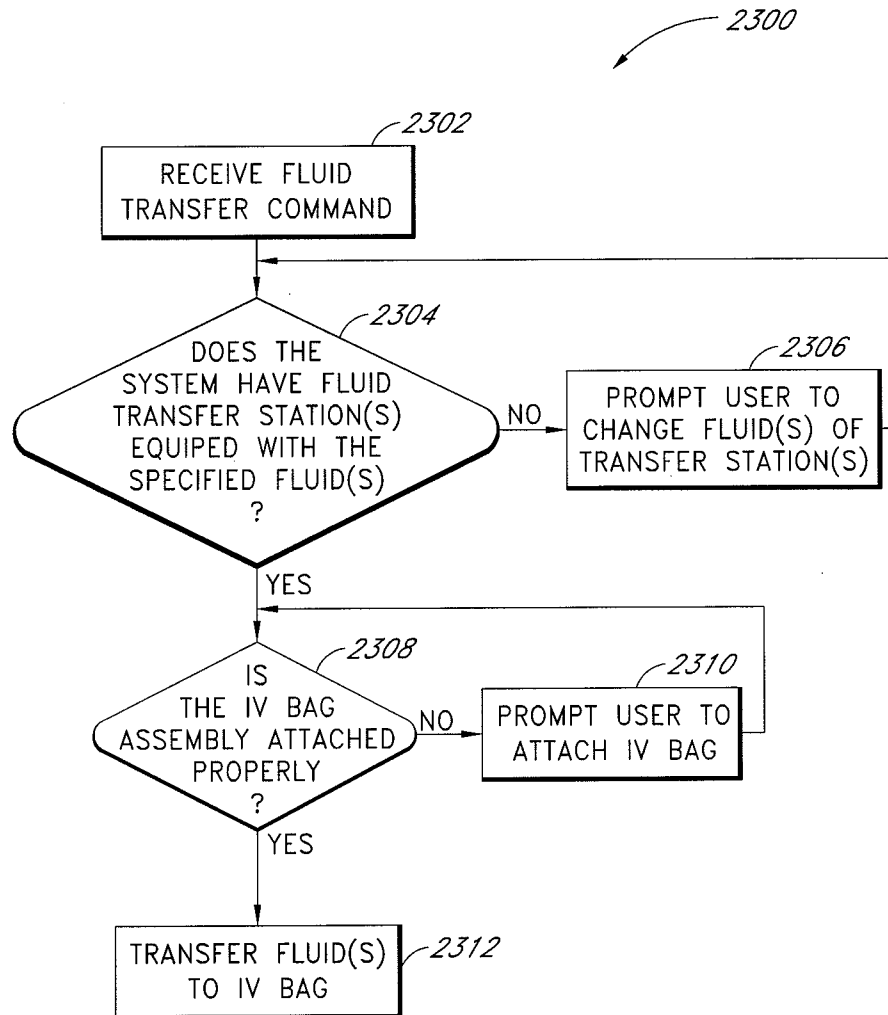
*FIG. 22*

FIG. 22A



*FIG. 23*

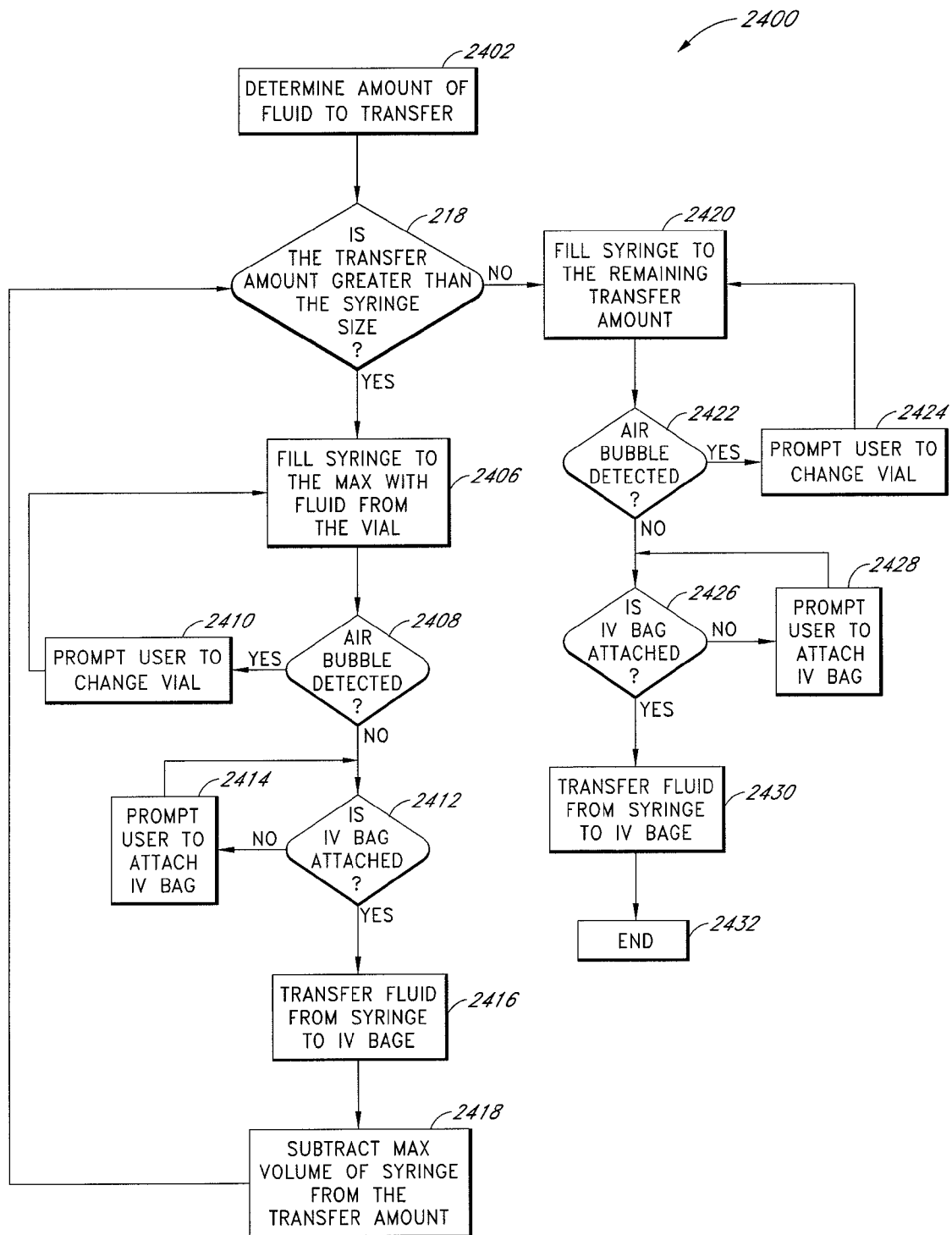
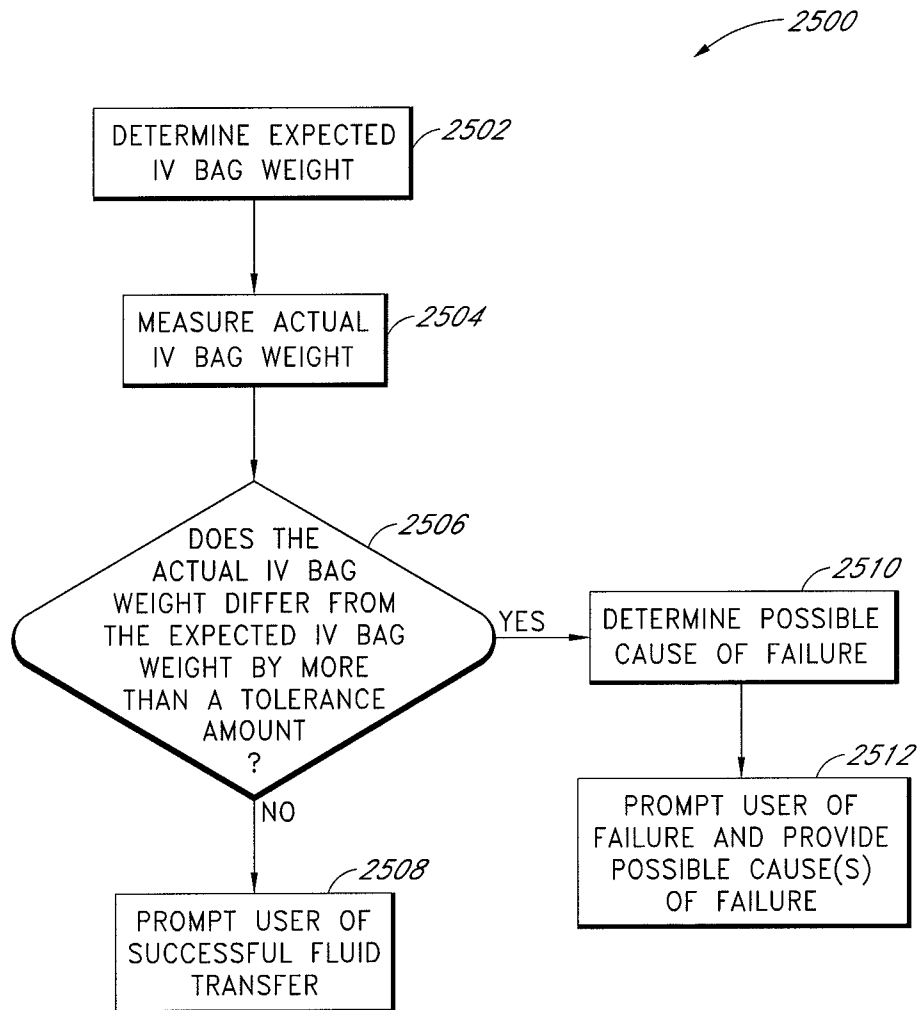


FIG. 24

*FIG. 25*

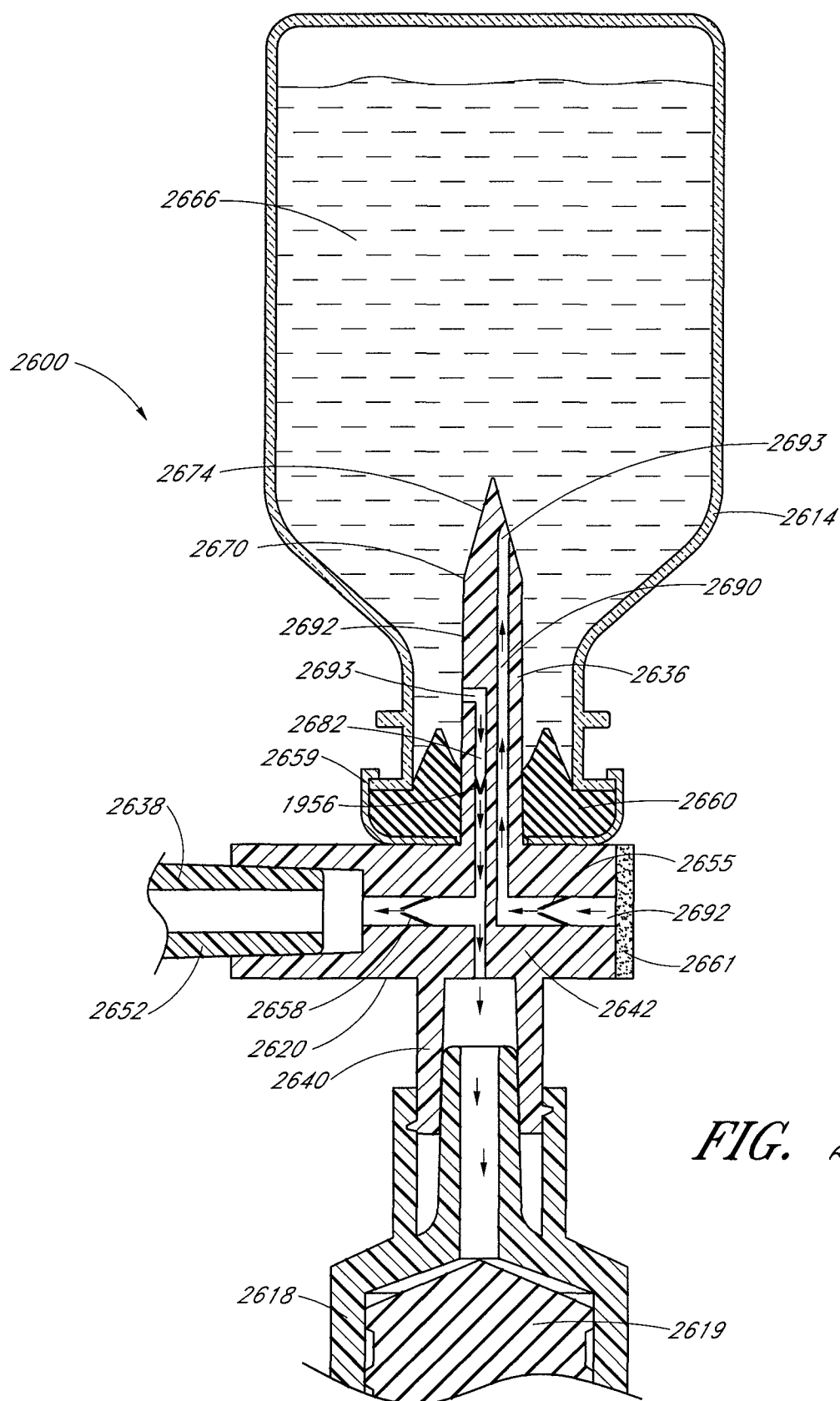
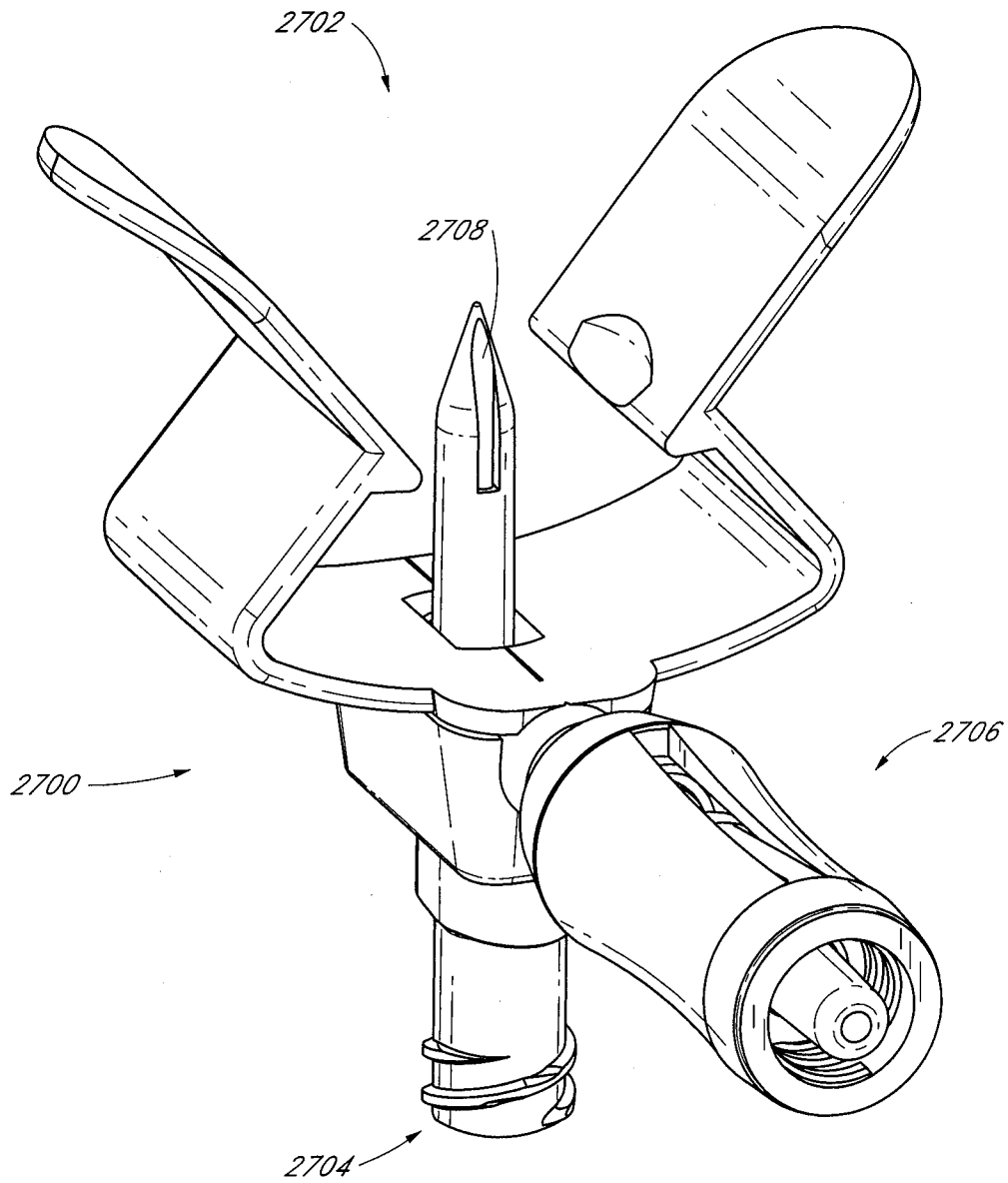
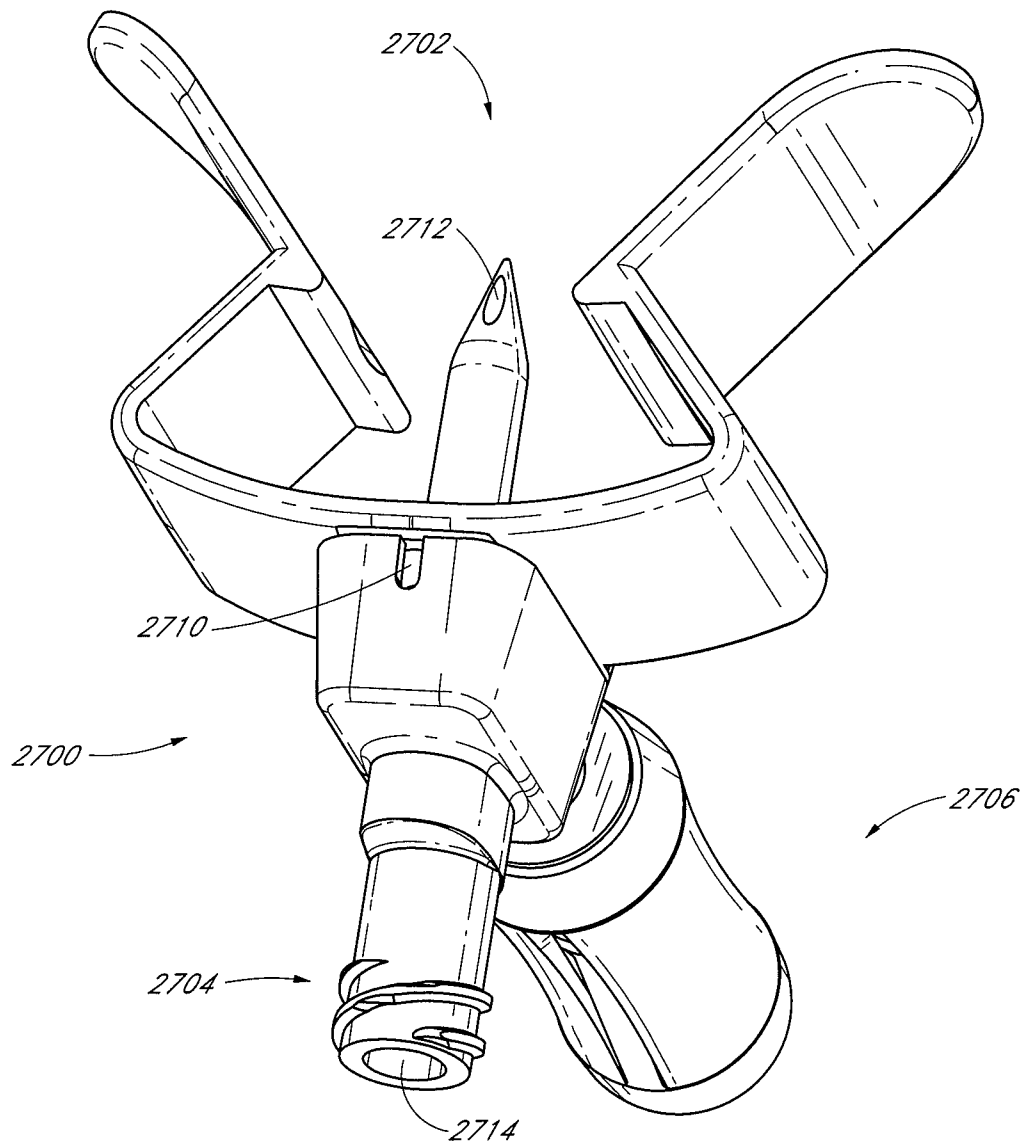


FIG. 26

*FIG. 27A*

*FIG. 27B*

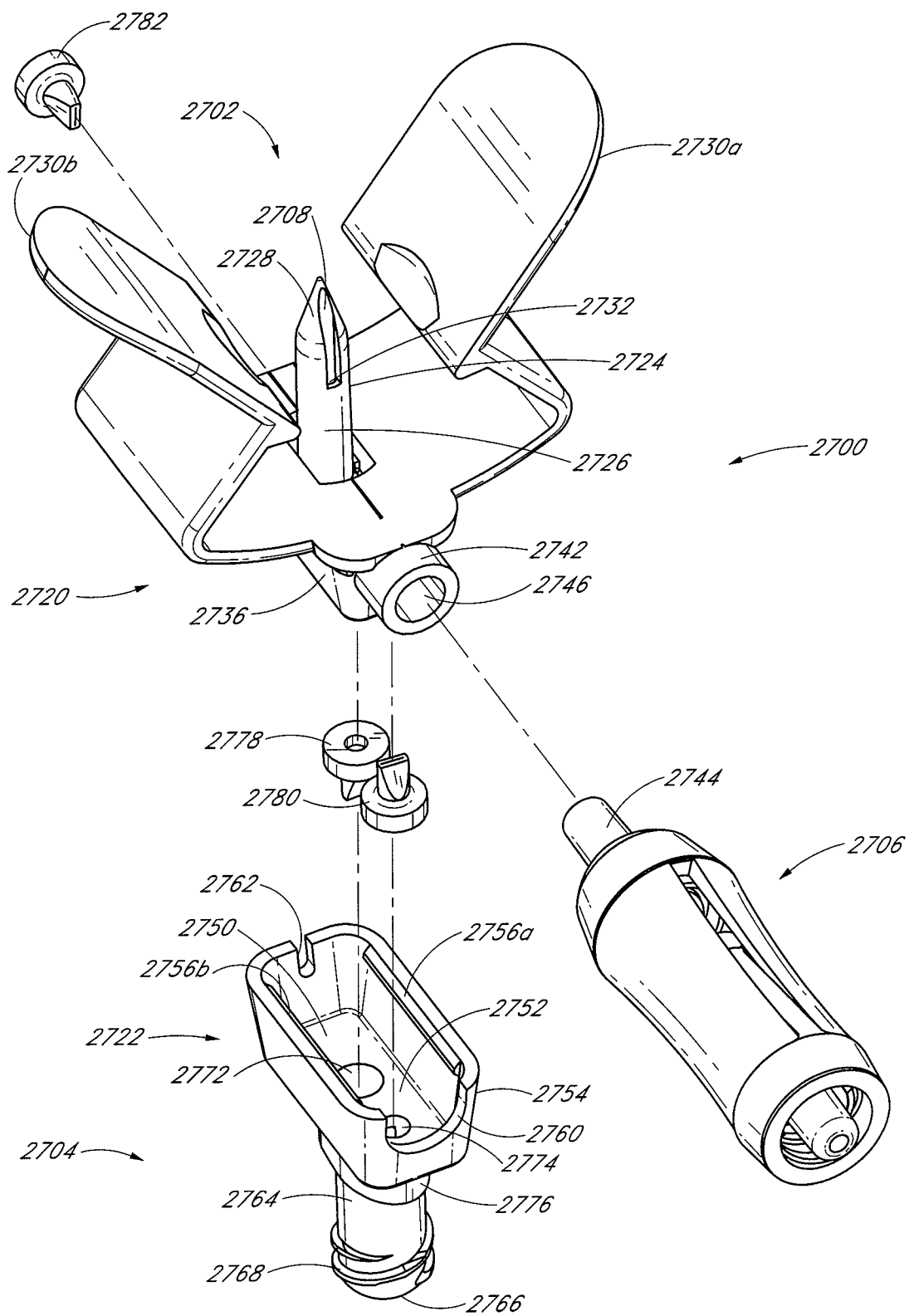


FIG. 28A

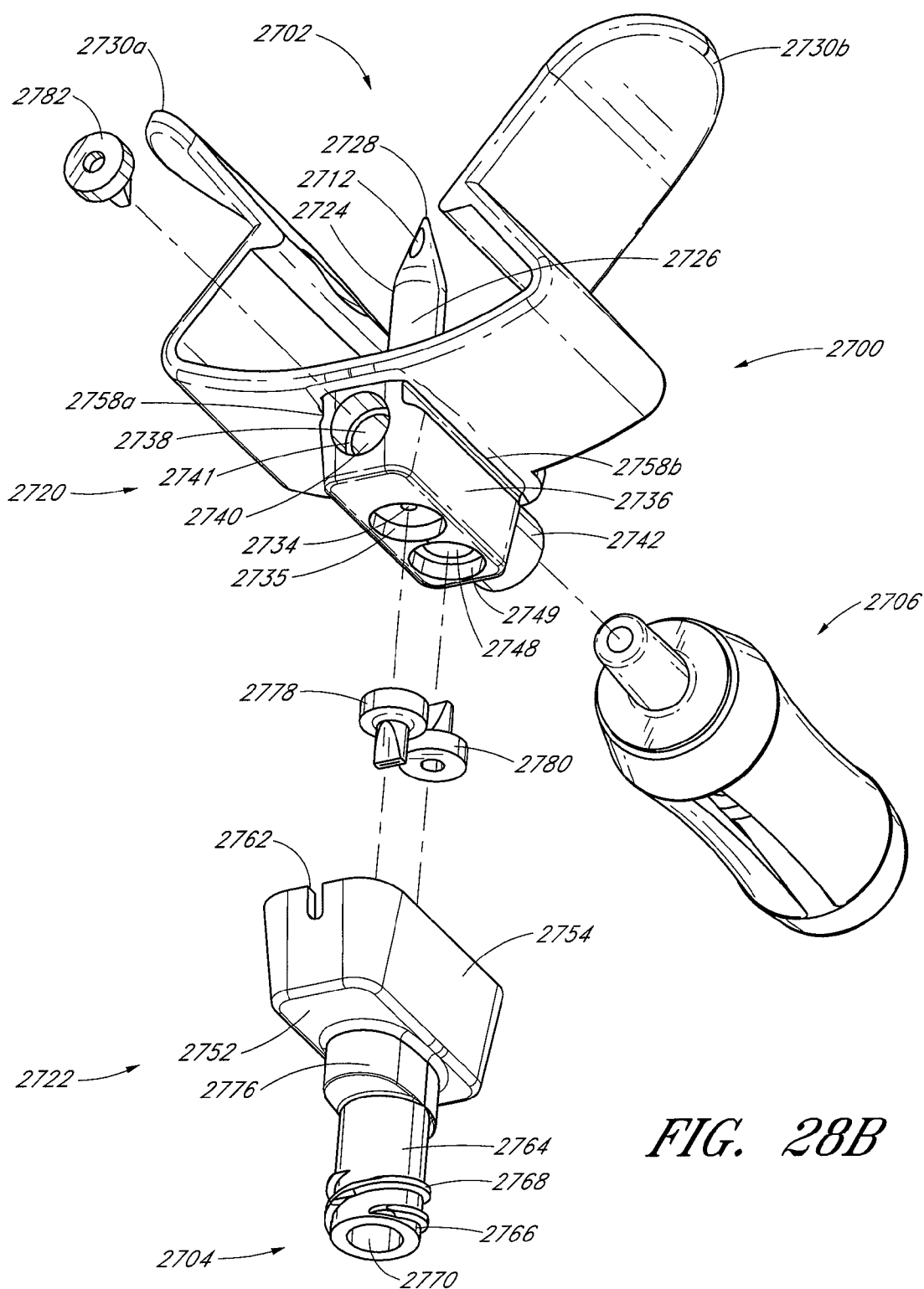
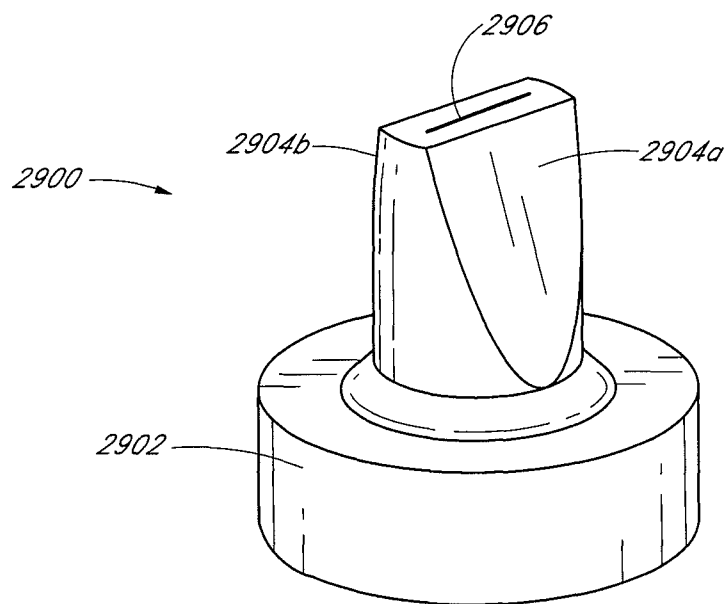
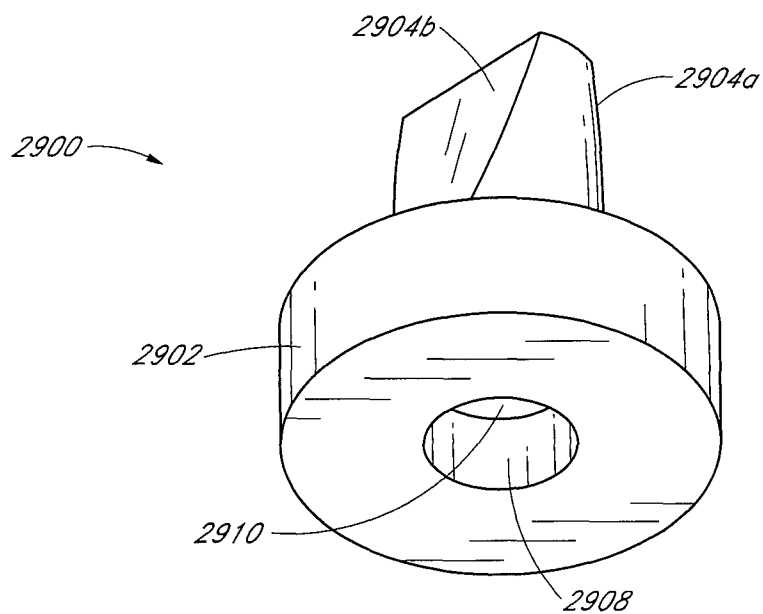


FIG. 28B

*FIG. 29A**FIG. 29B*

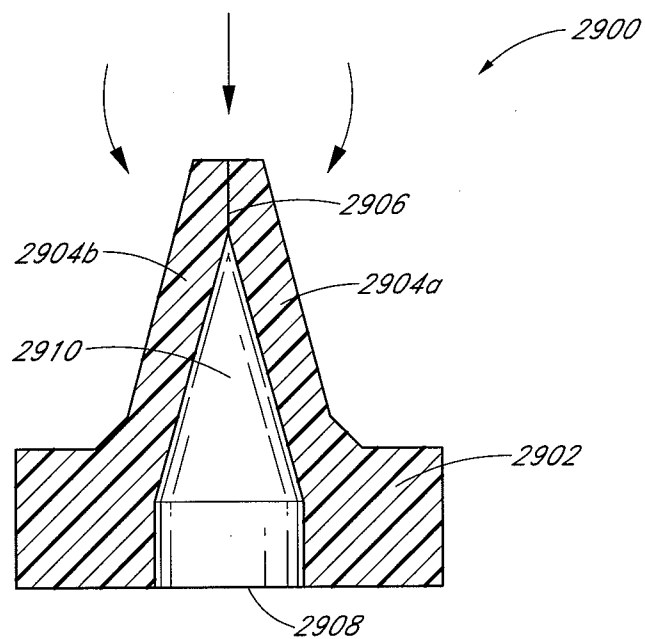


FIG. 29C

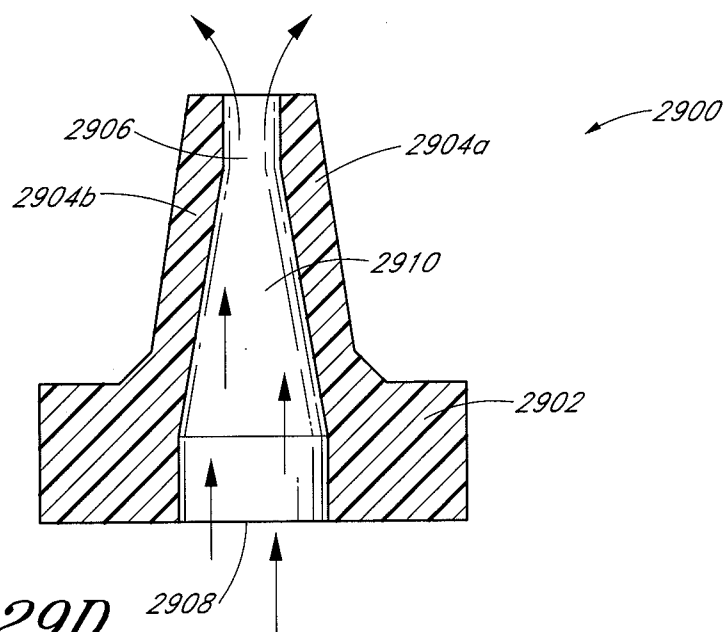
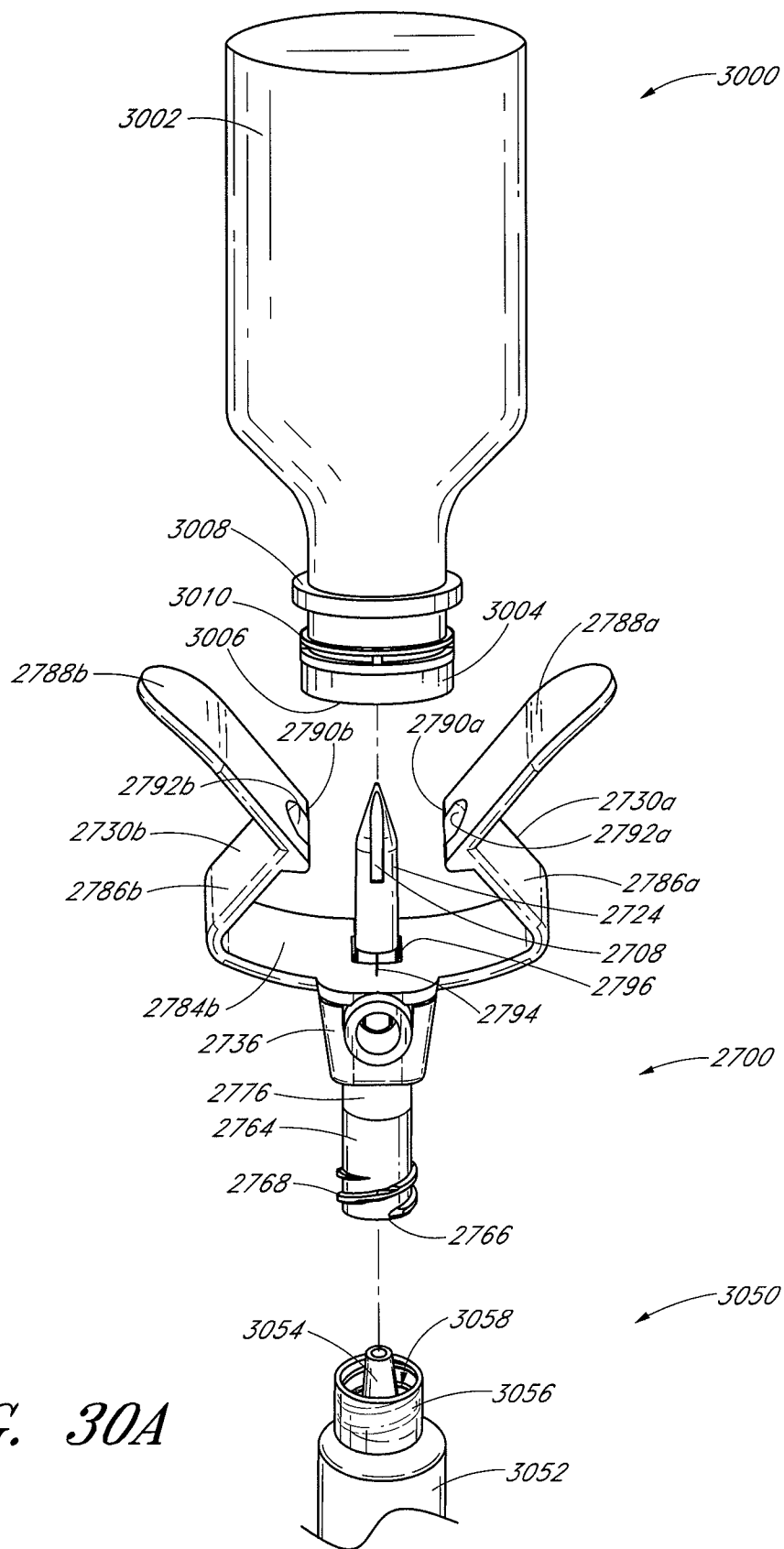


FIG. 29D



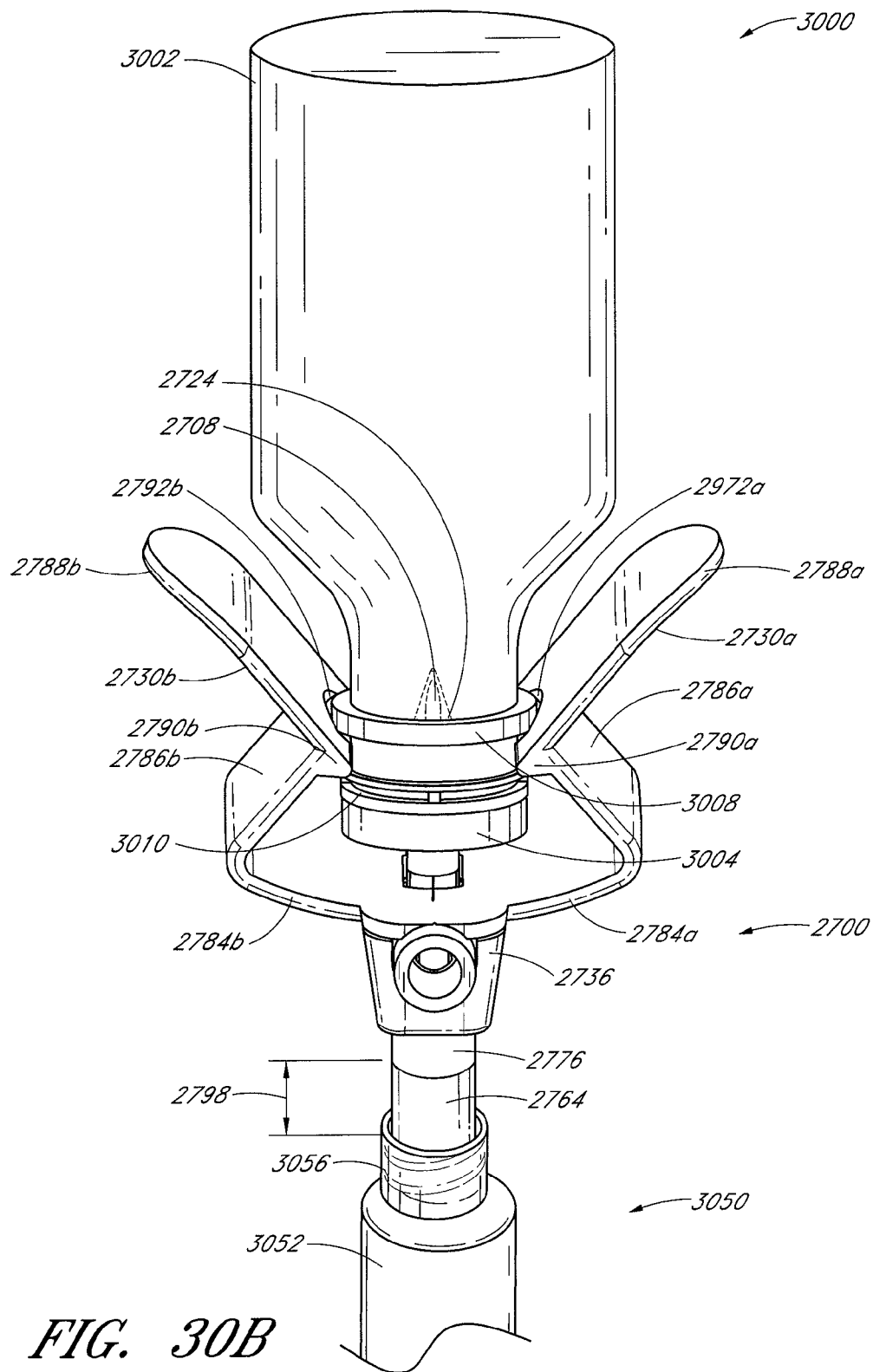
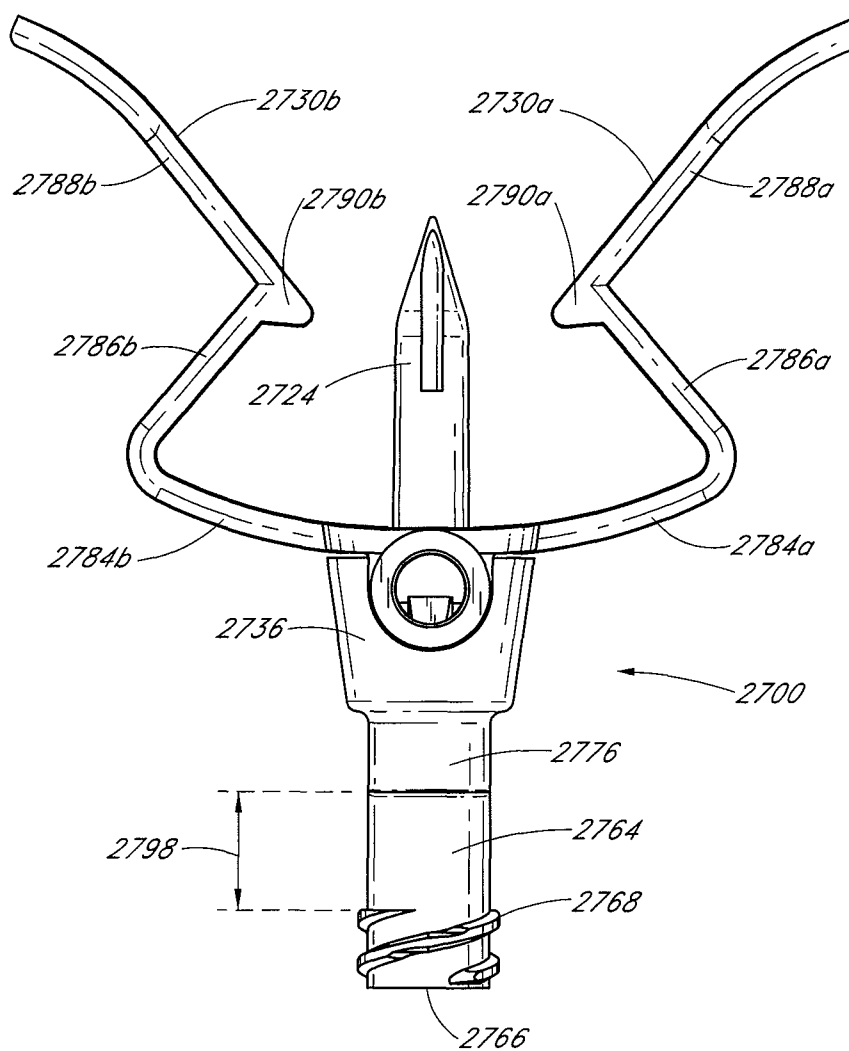


FIG. 30B

*FIG. 30C*

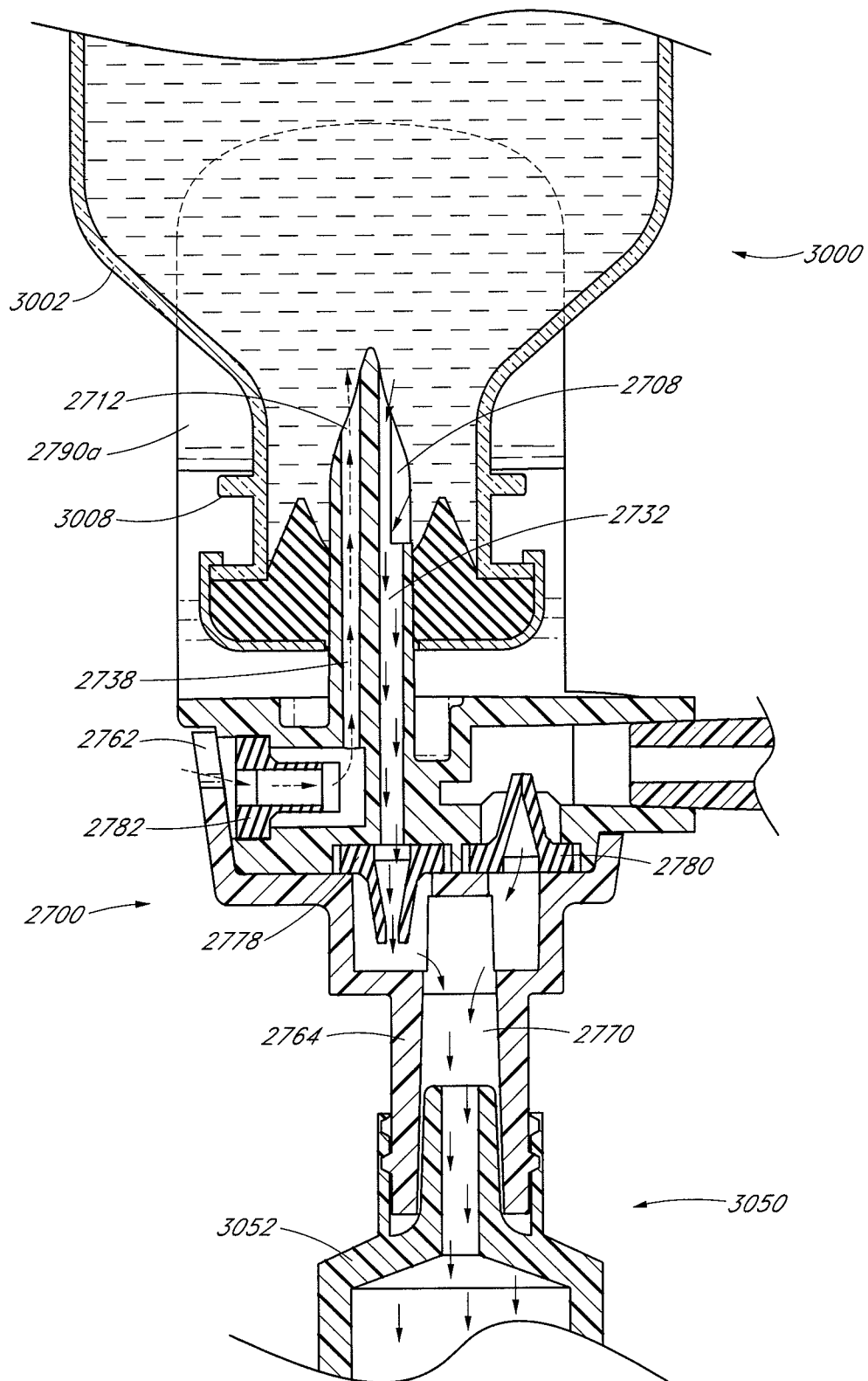


FIG. 31A

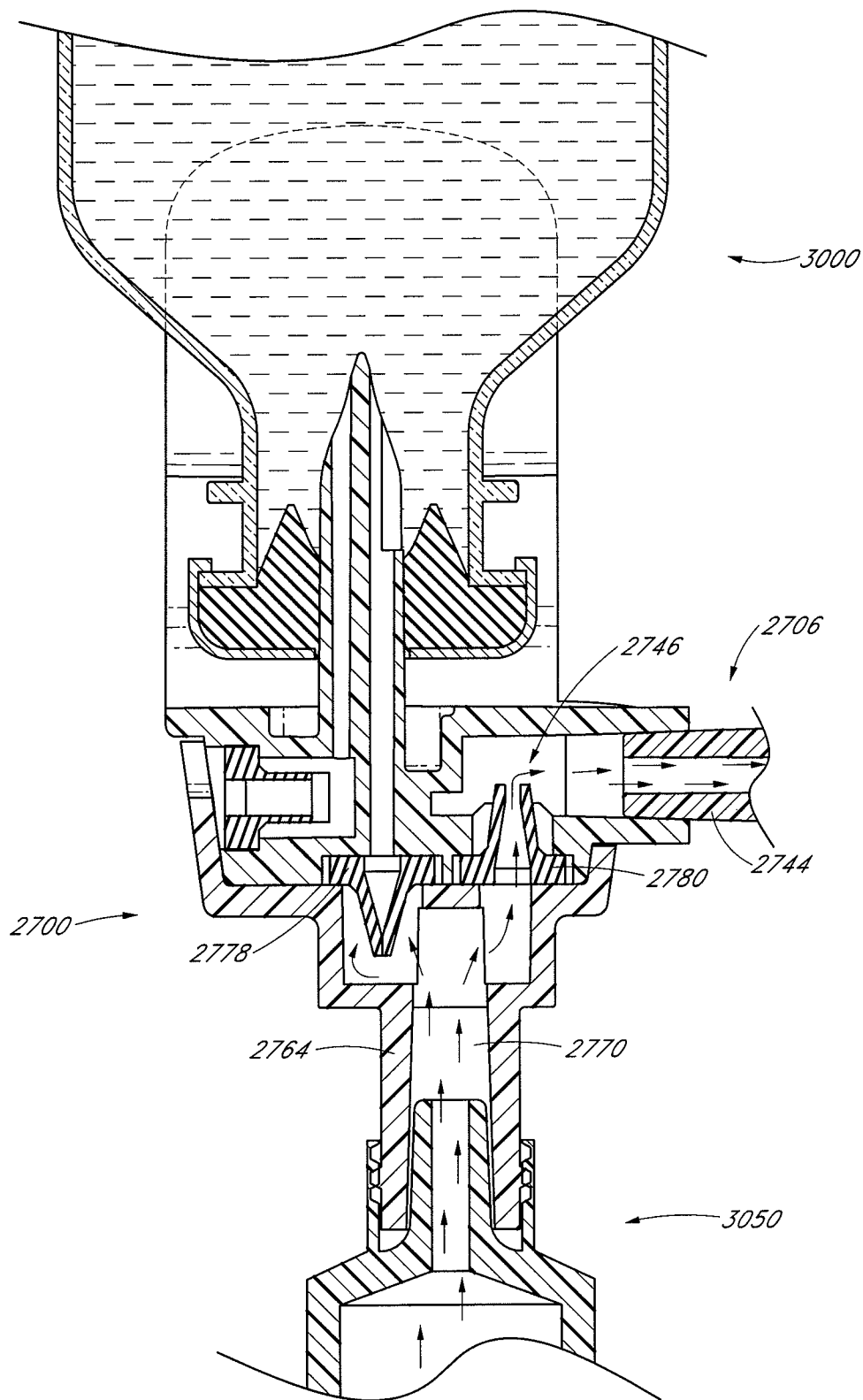


FIG. 31B

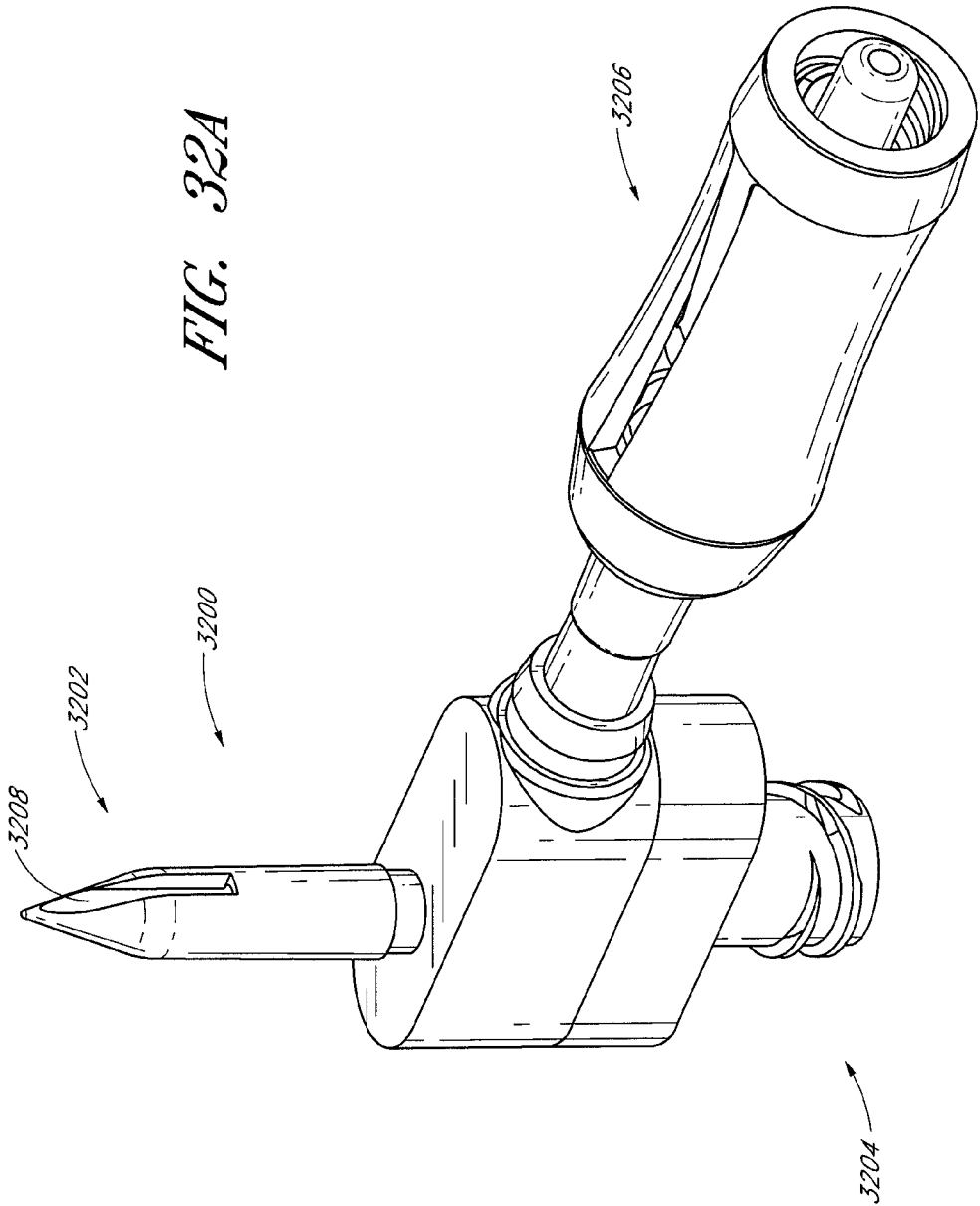
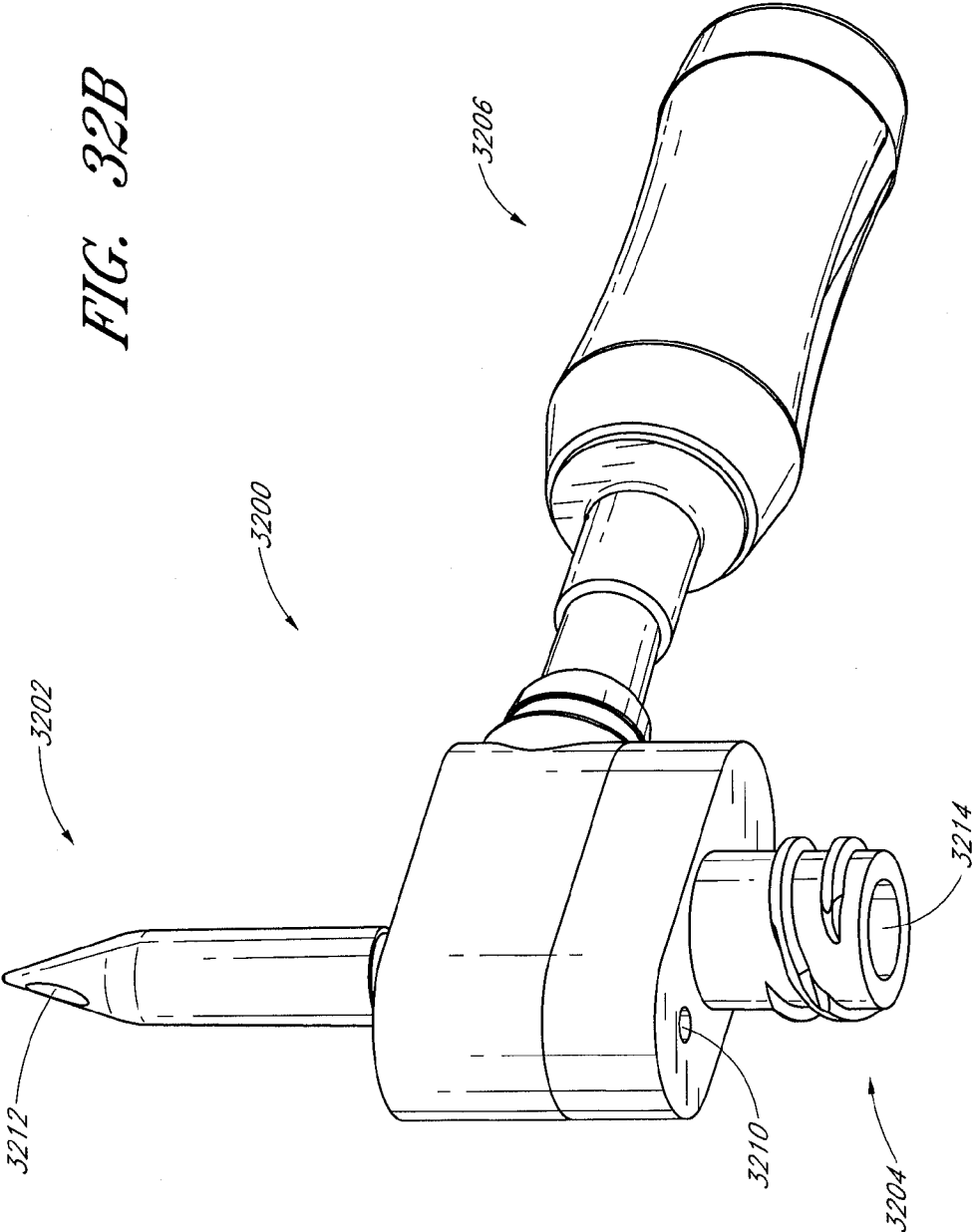


FIG. 32B



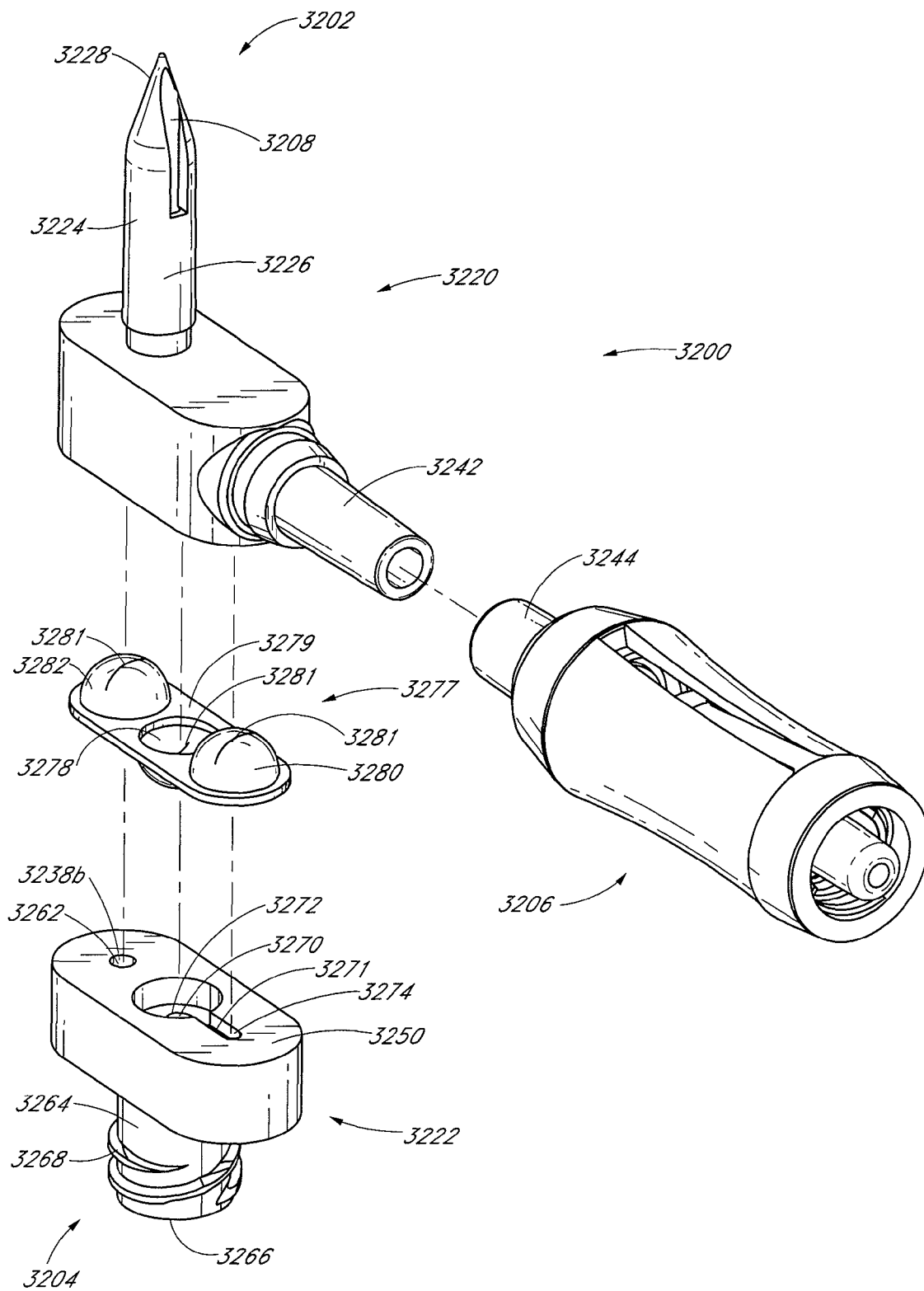


FIG. 33A

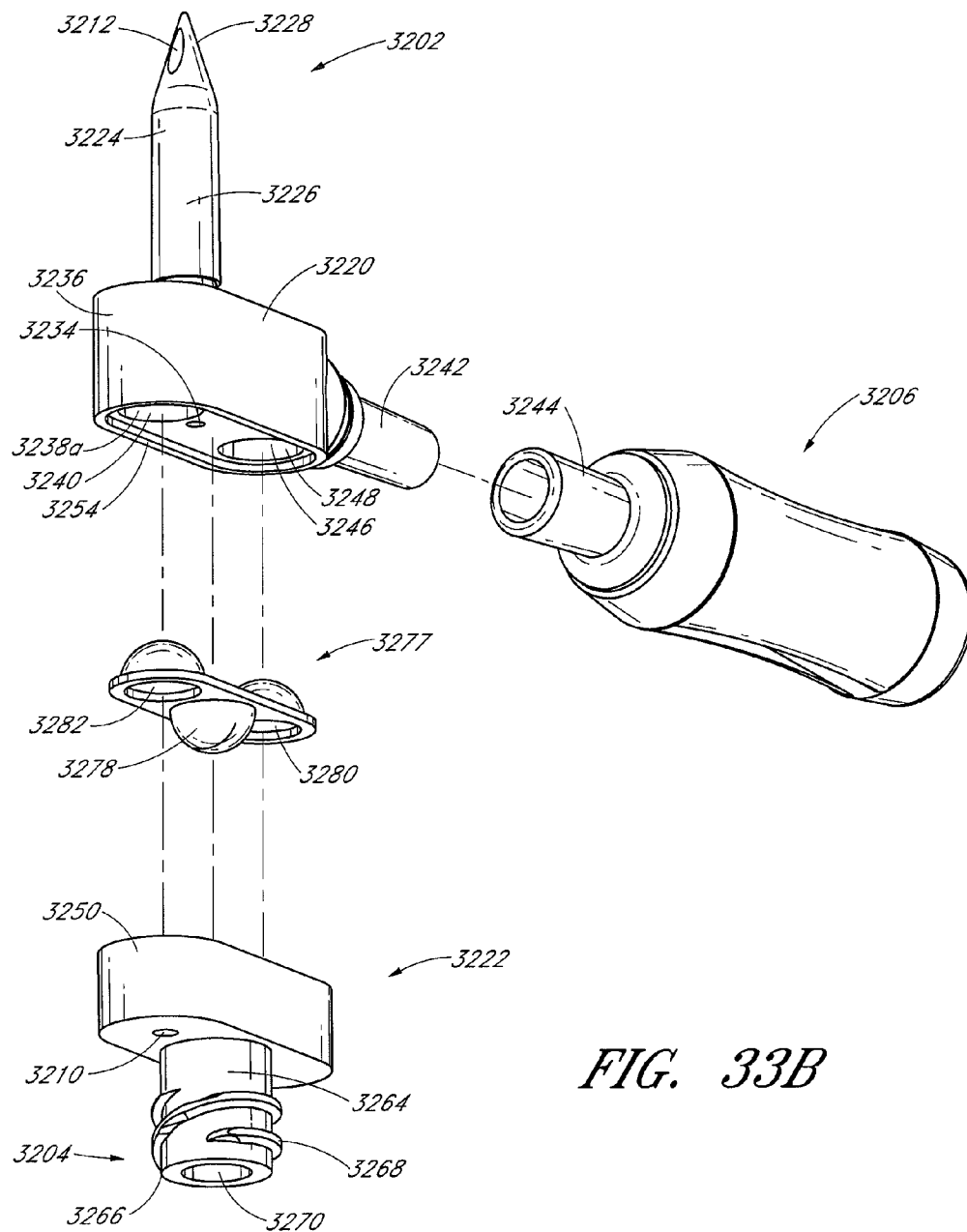


FIG. 33B

FIG. 34A

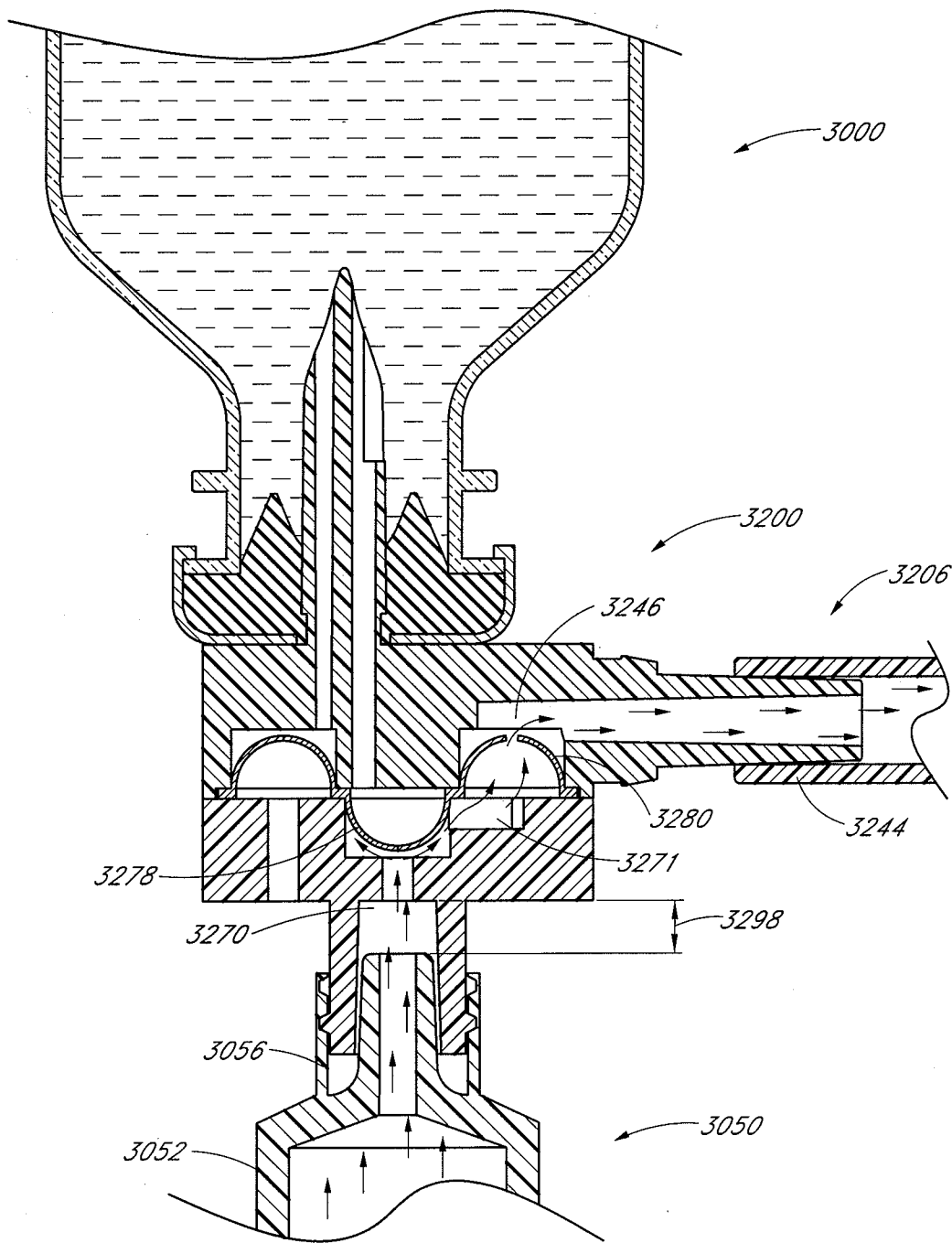
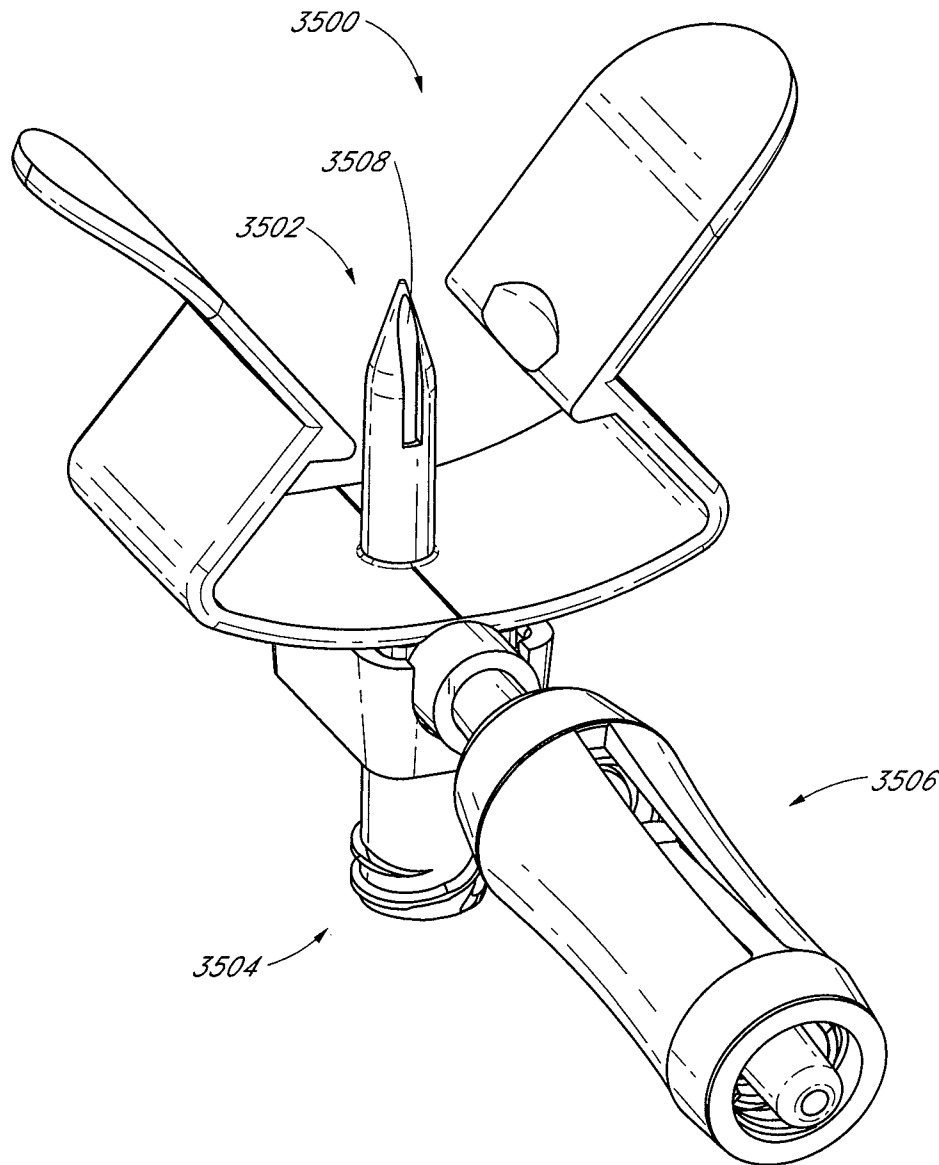
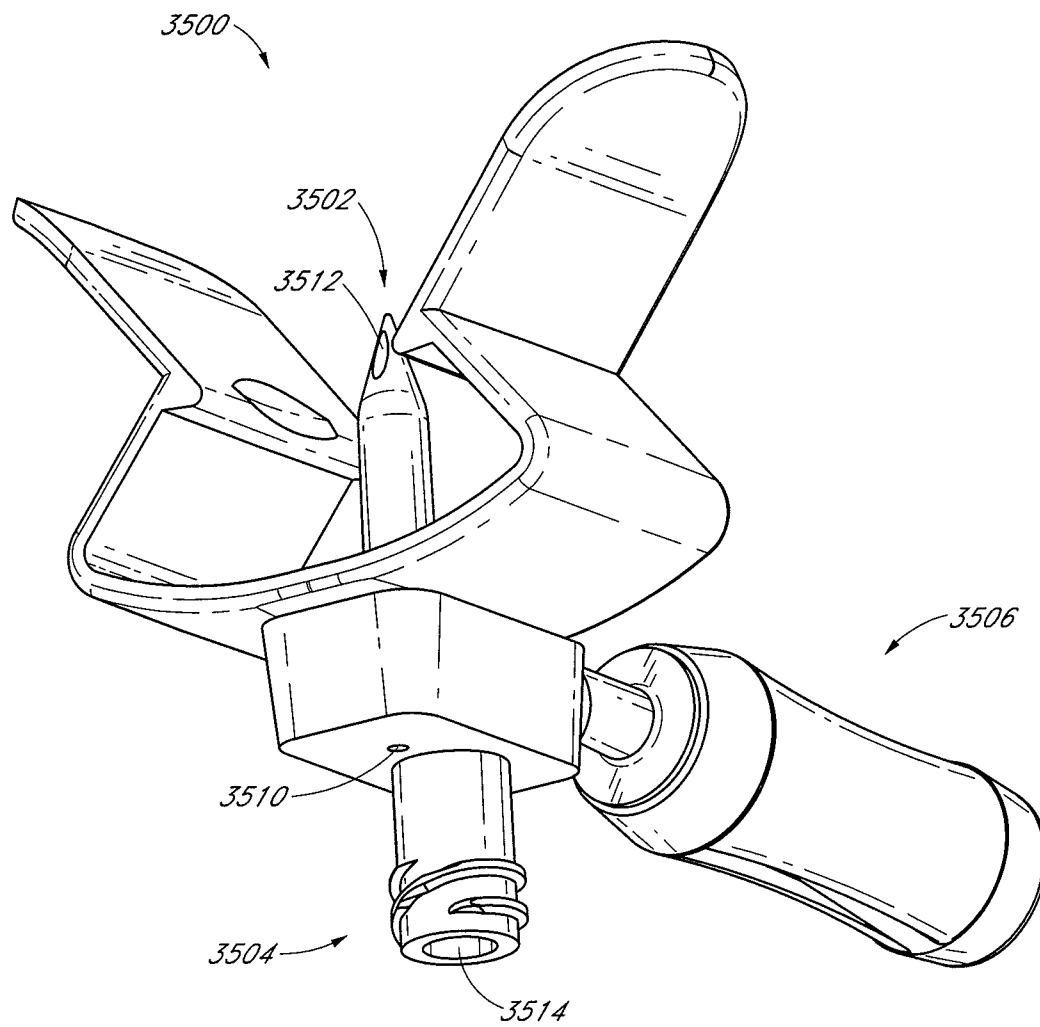
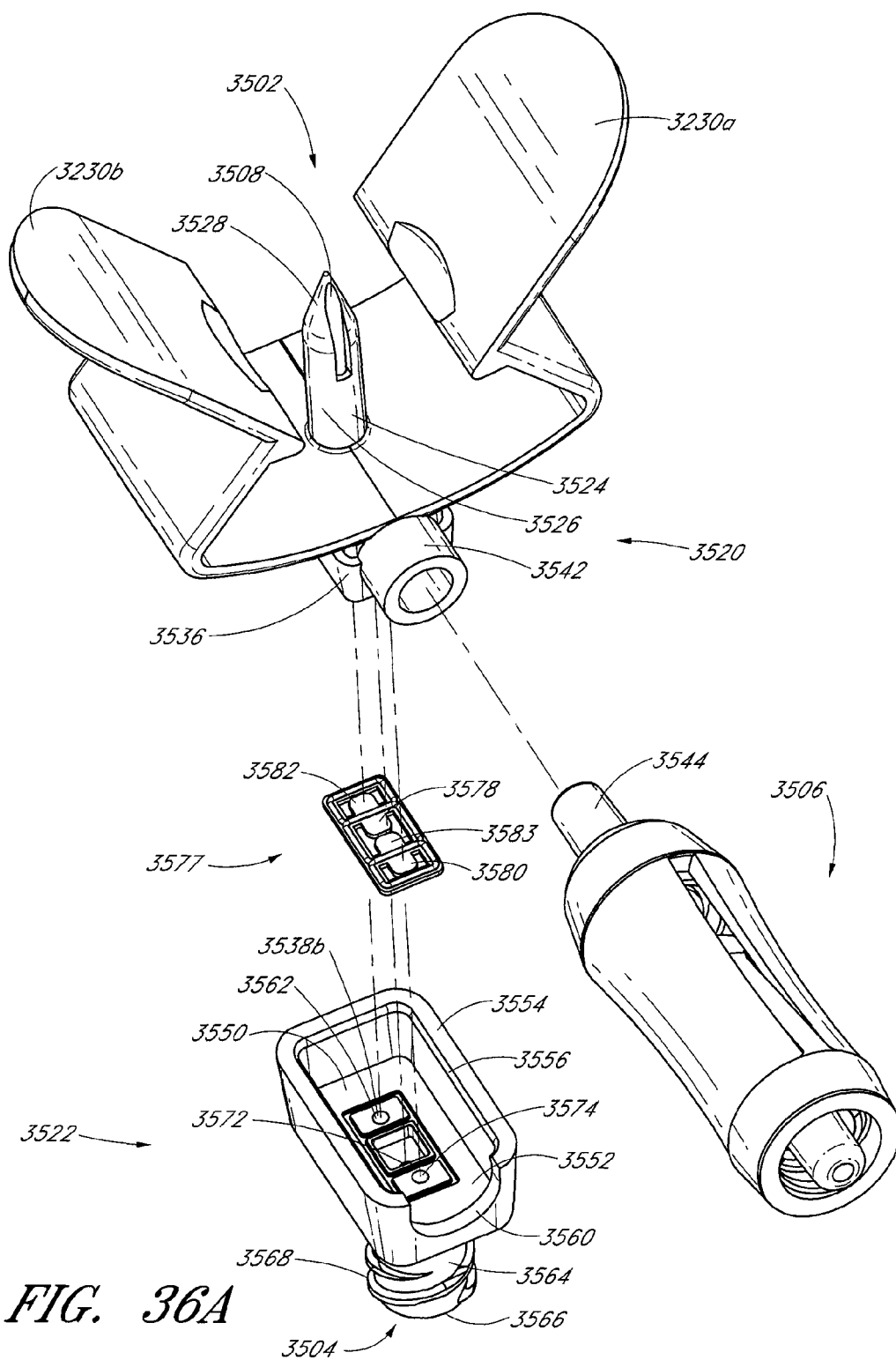


FIG. 34B

*FIG. 35A*

*FIG. 35B*



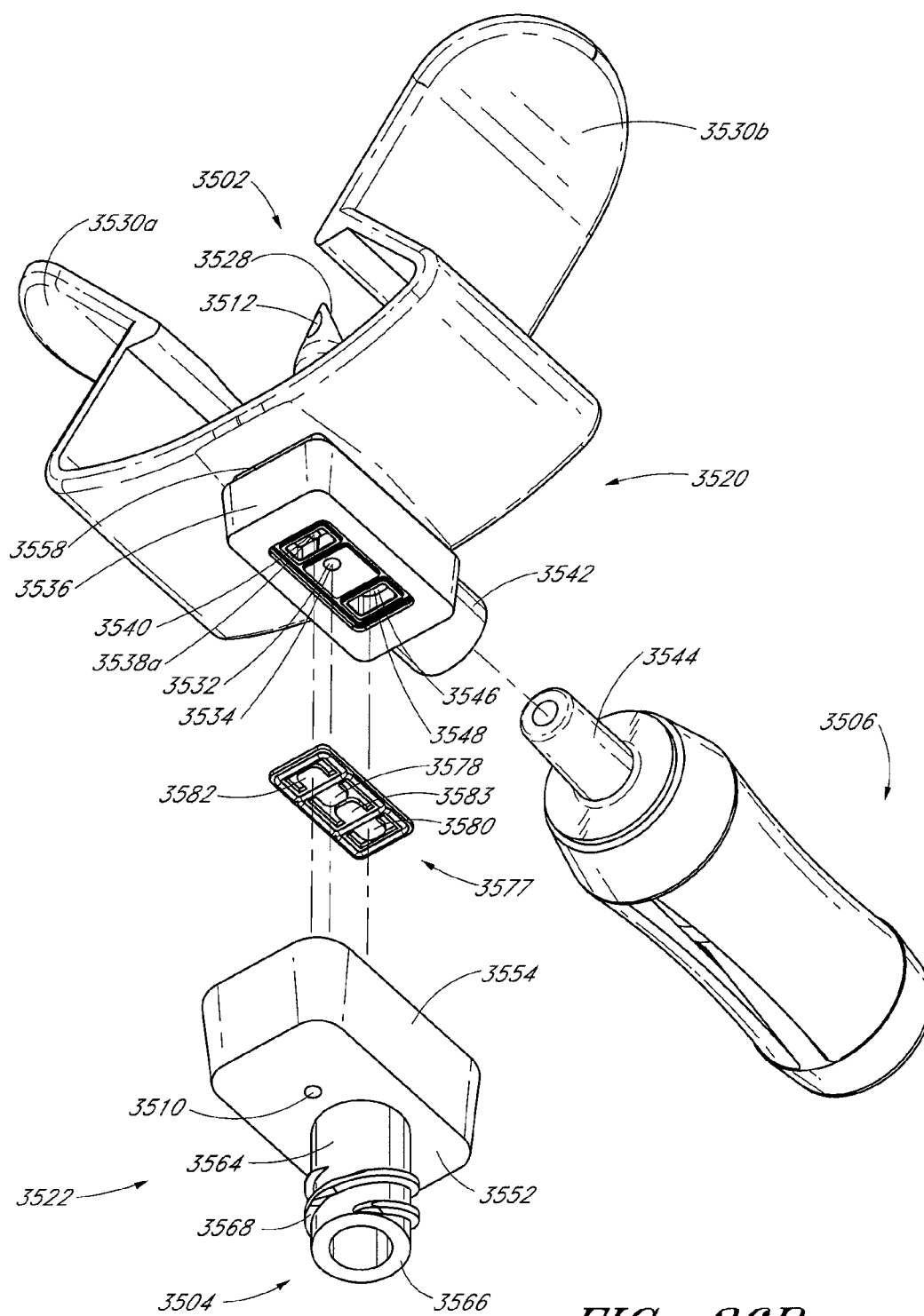


FIG. 36B

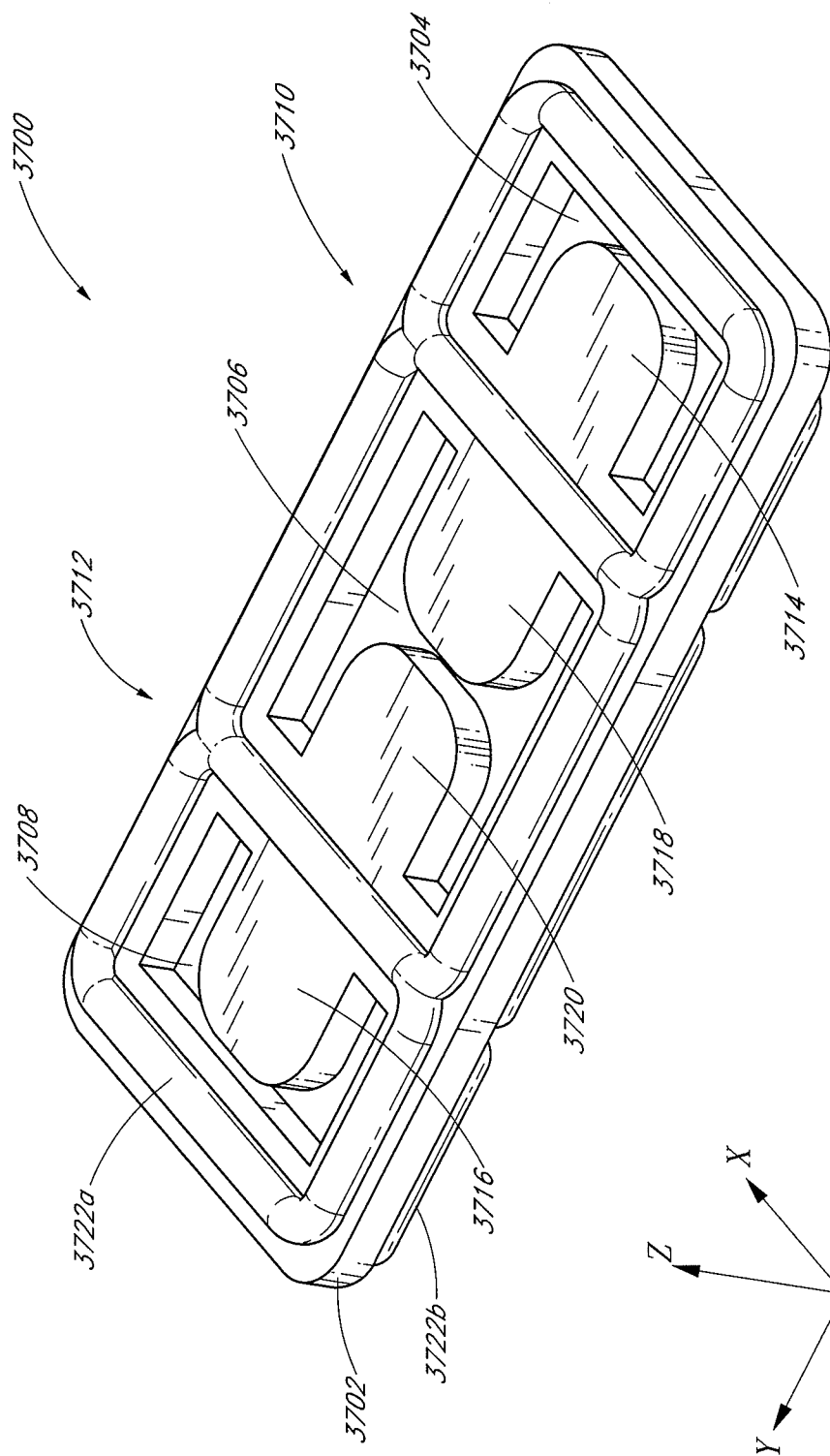


FIG. 37

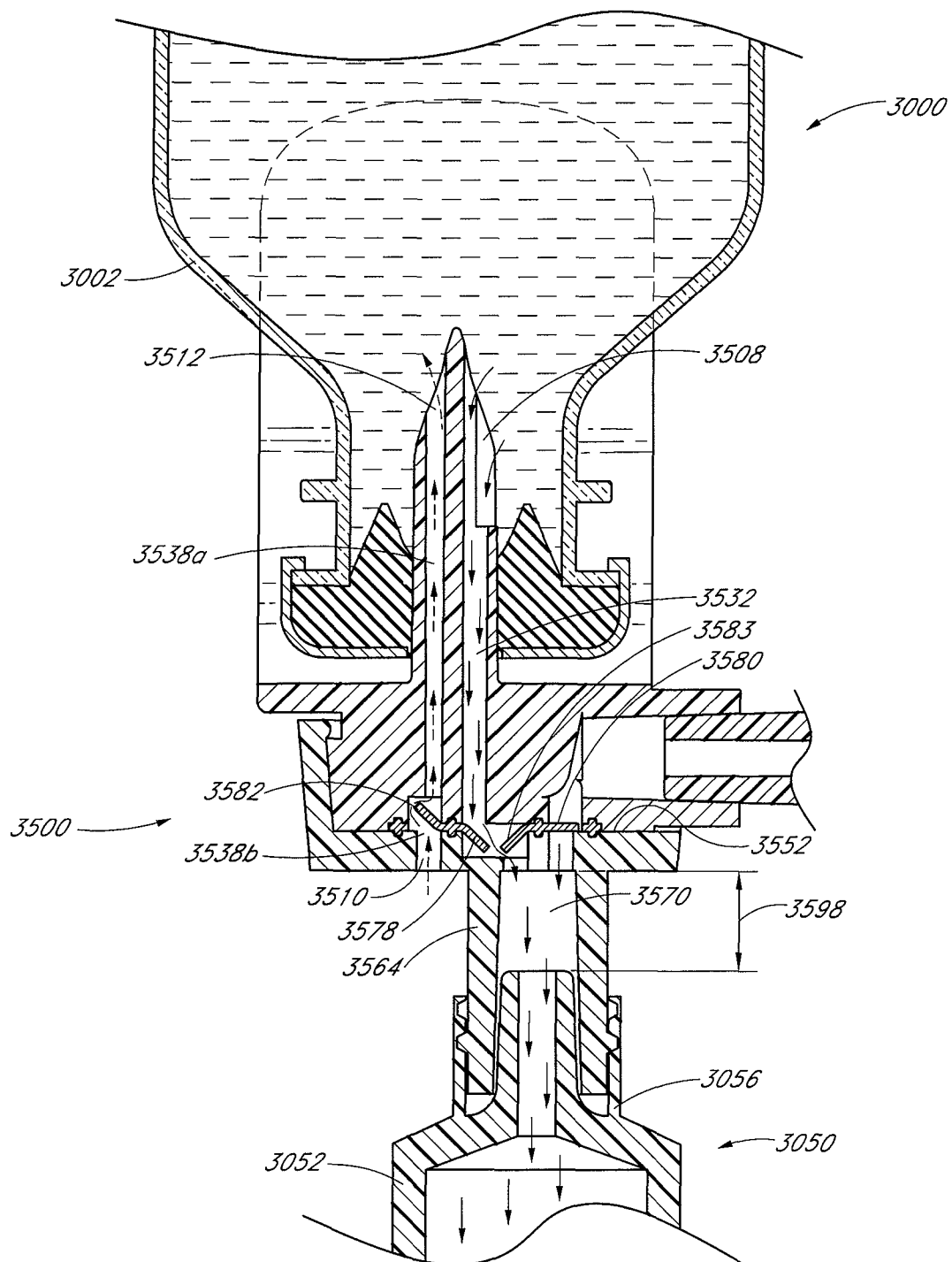


FIG. 38A

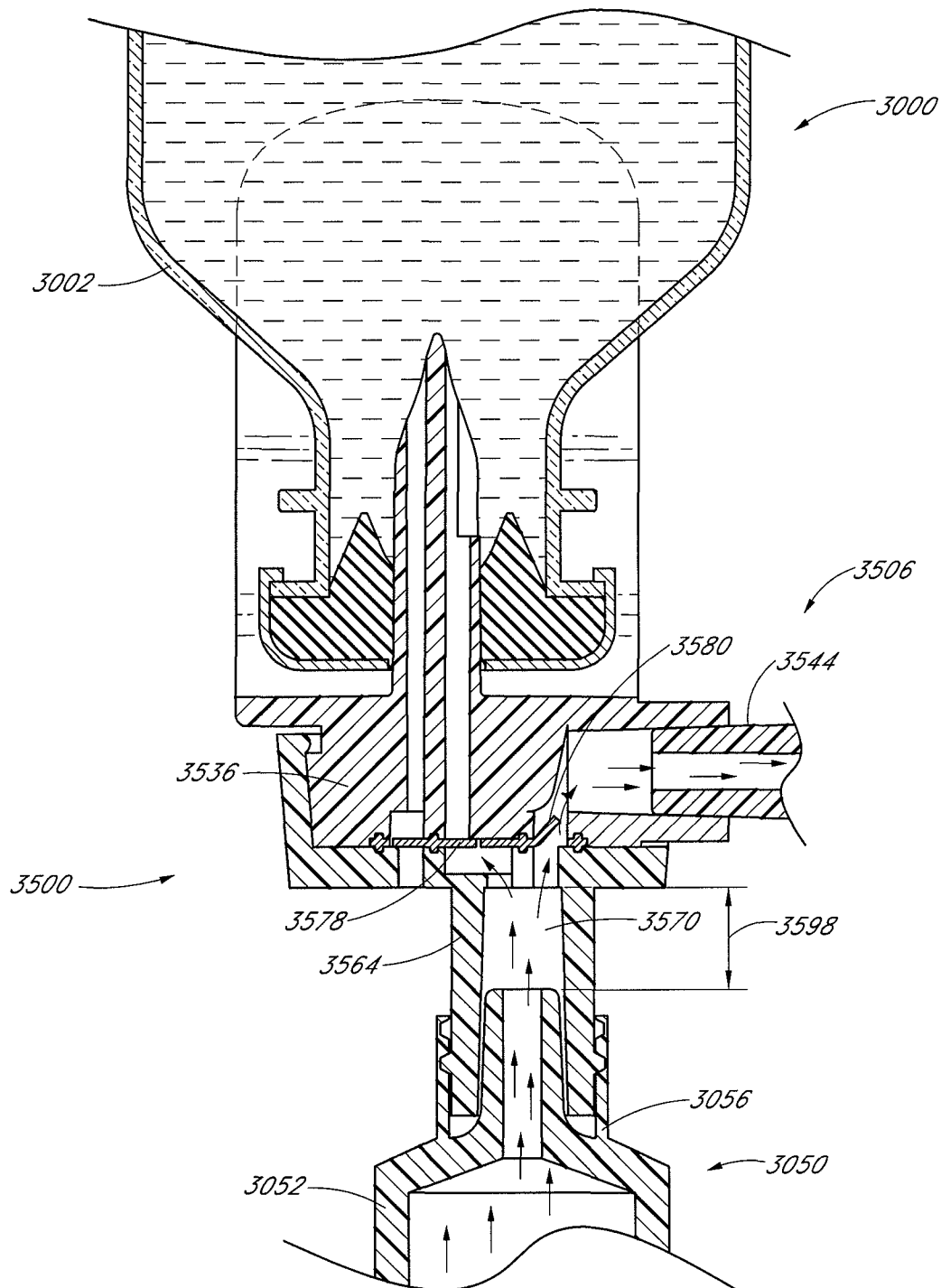
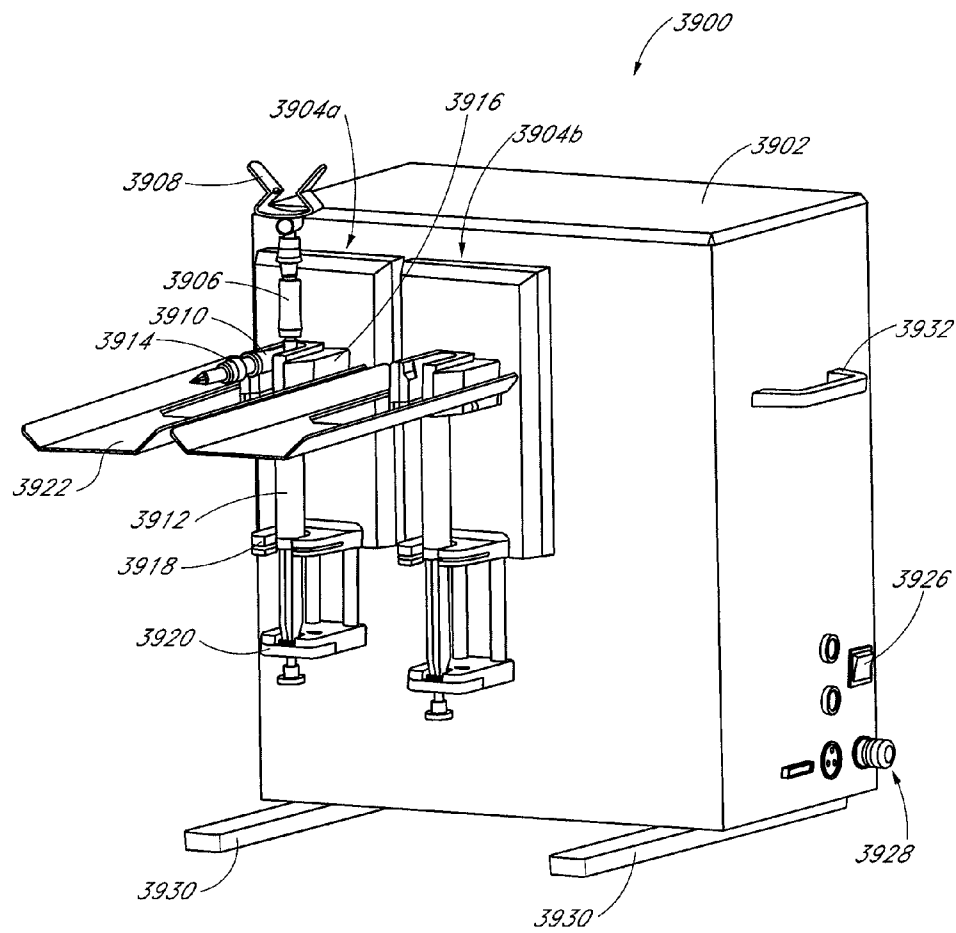


FIG. 38B

*FIG. 39*

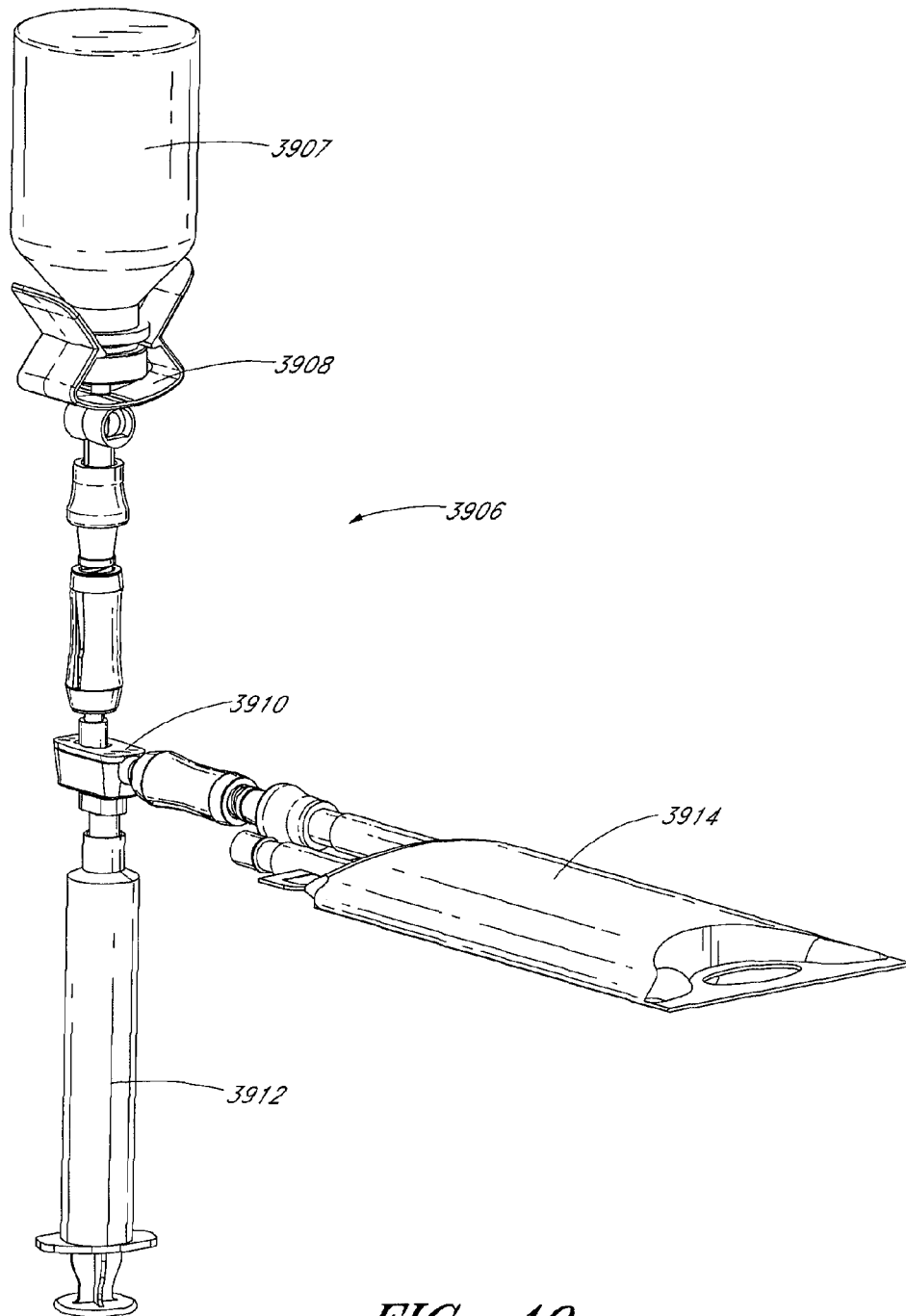


FIG. 40

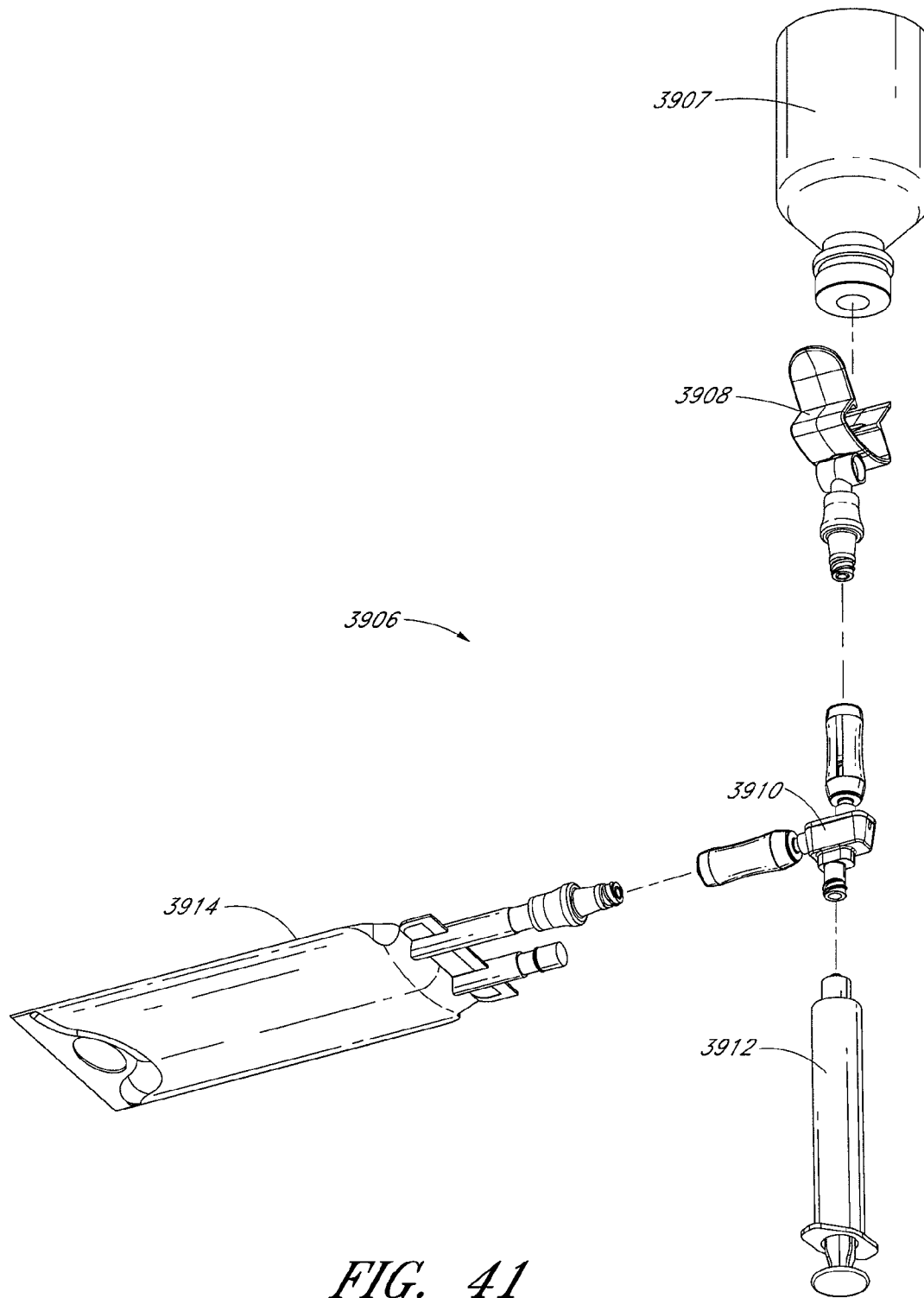


FIG. 41

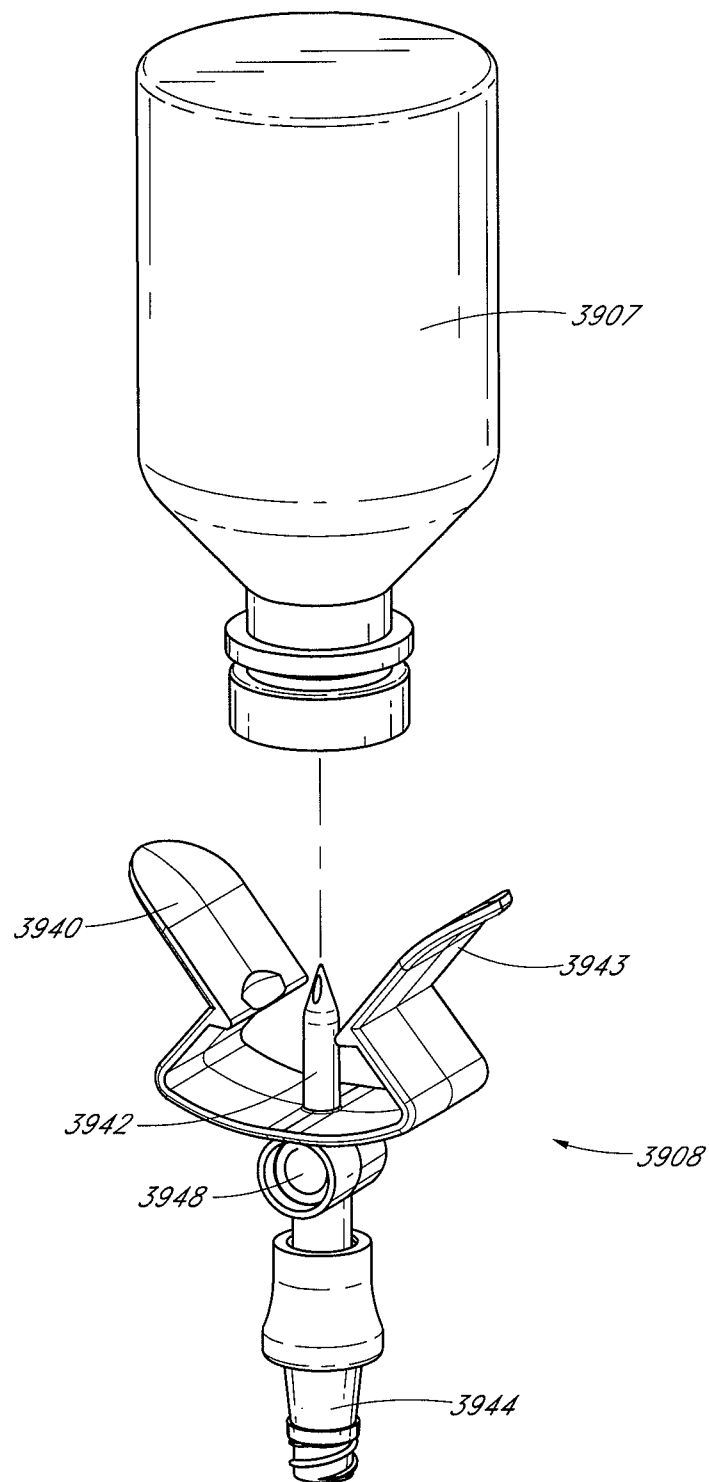


FIG. 42

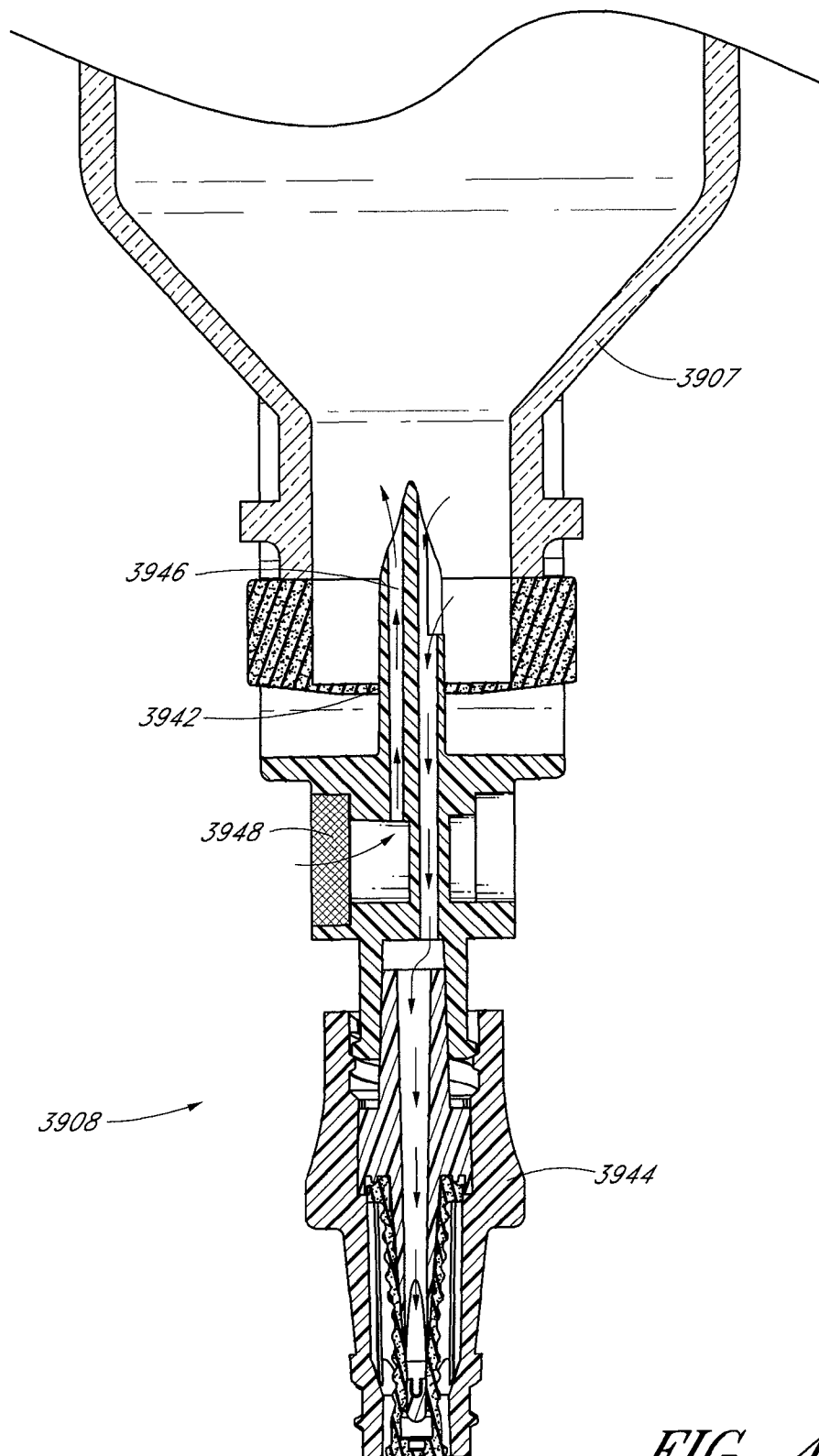


FIG. 43

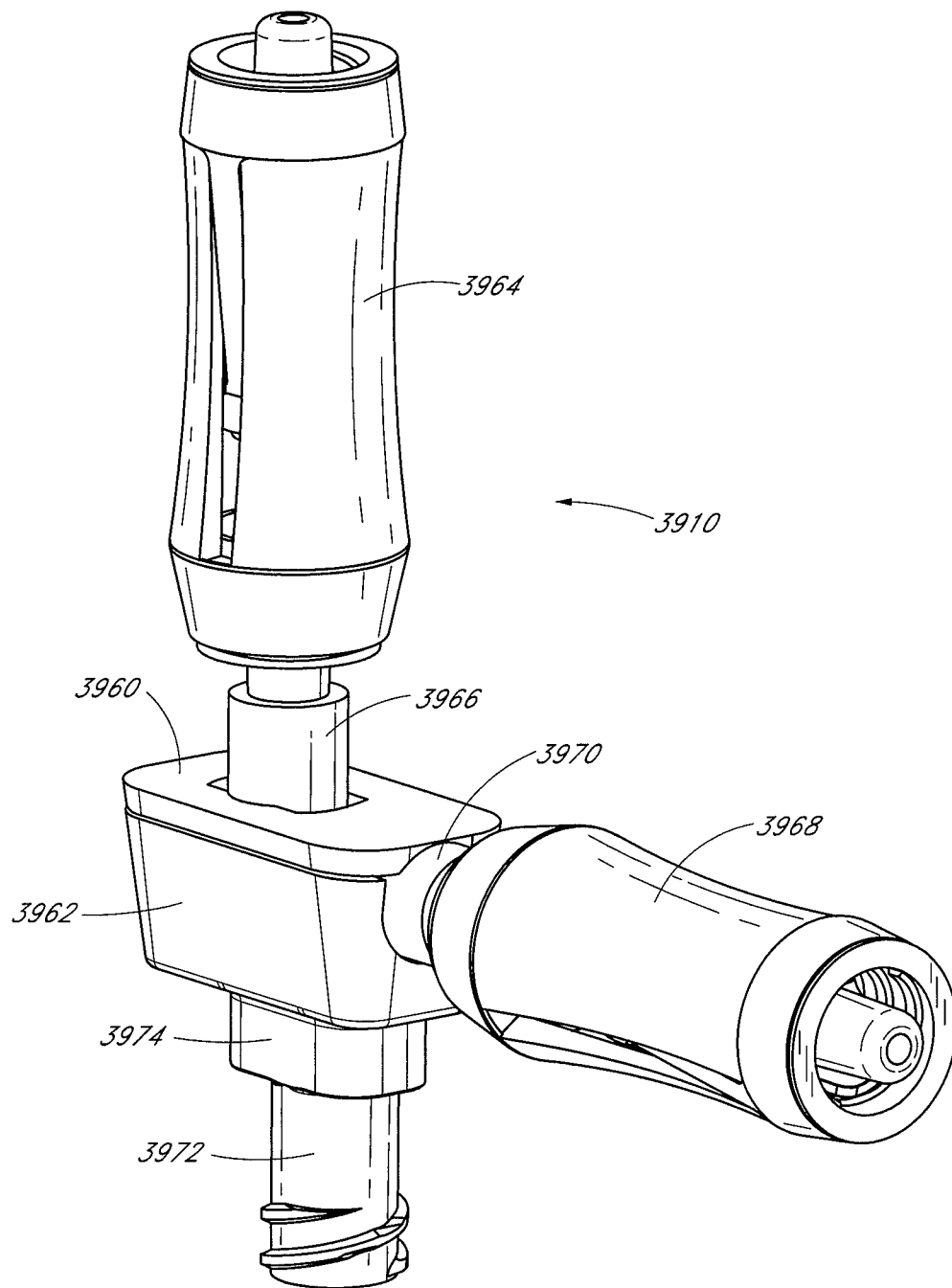
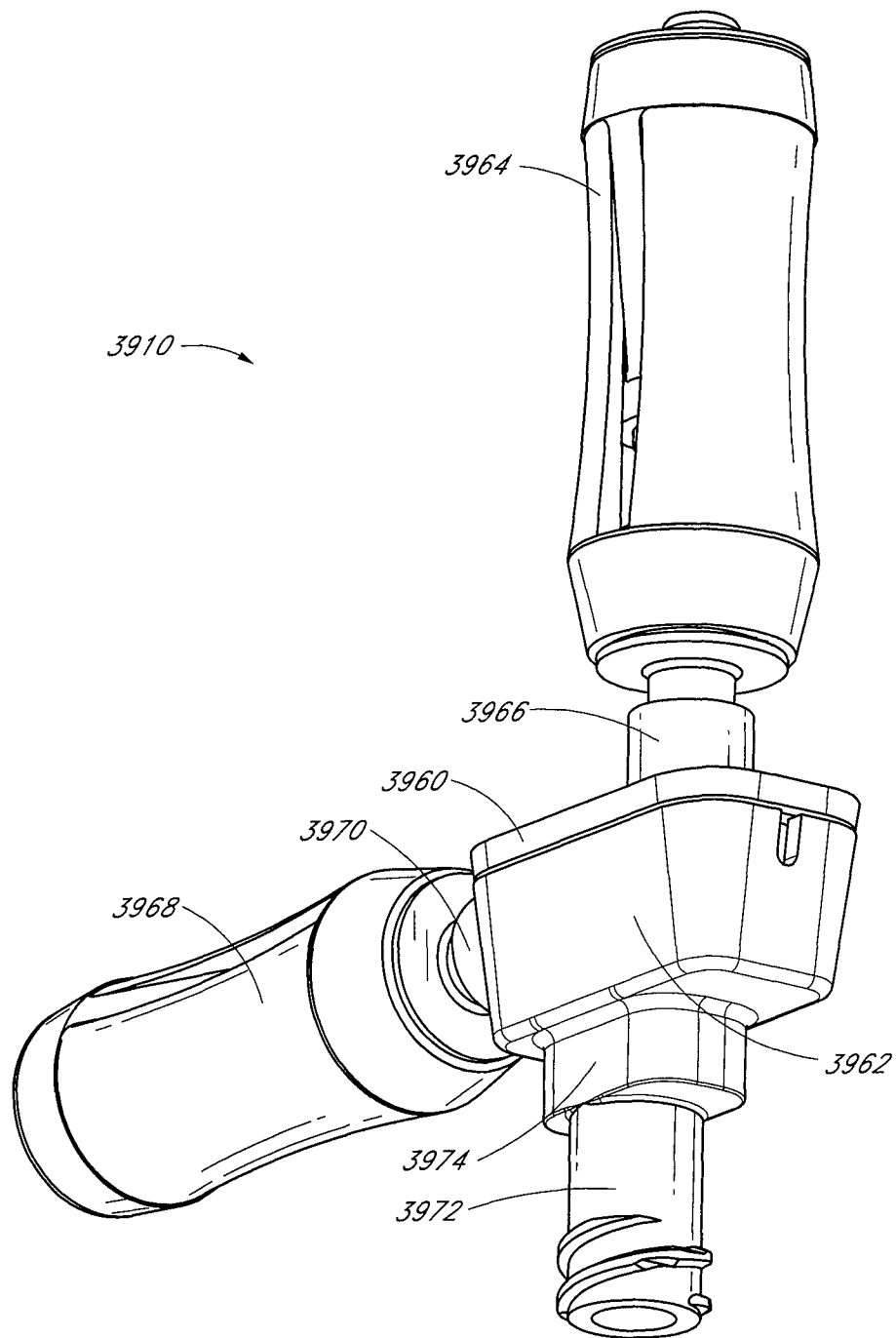
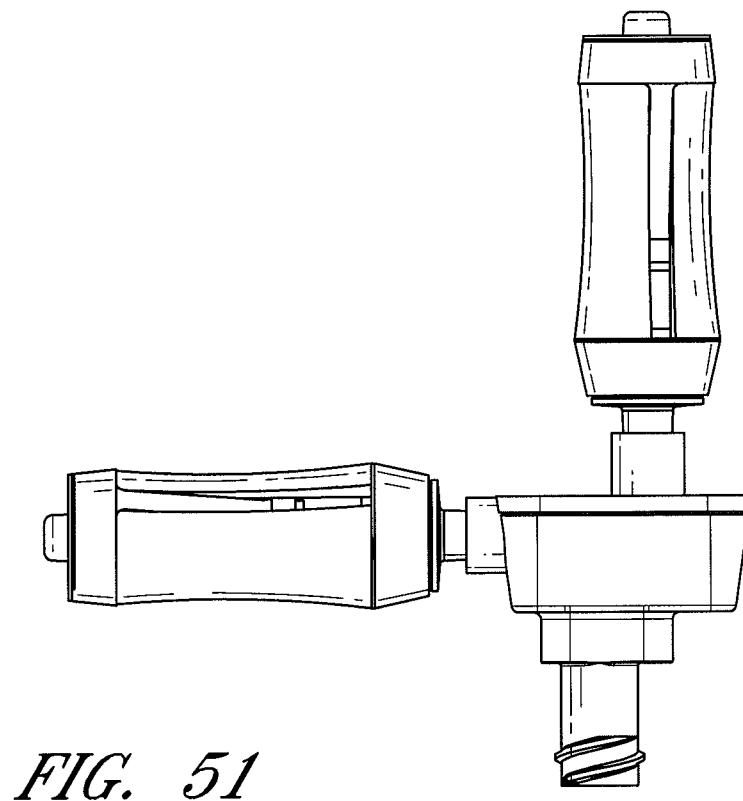
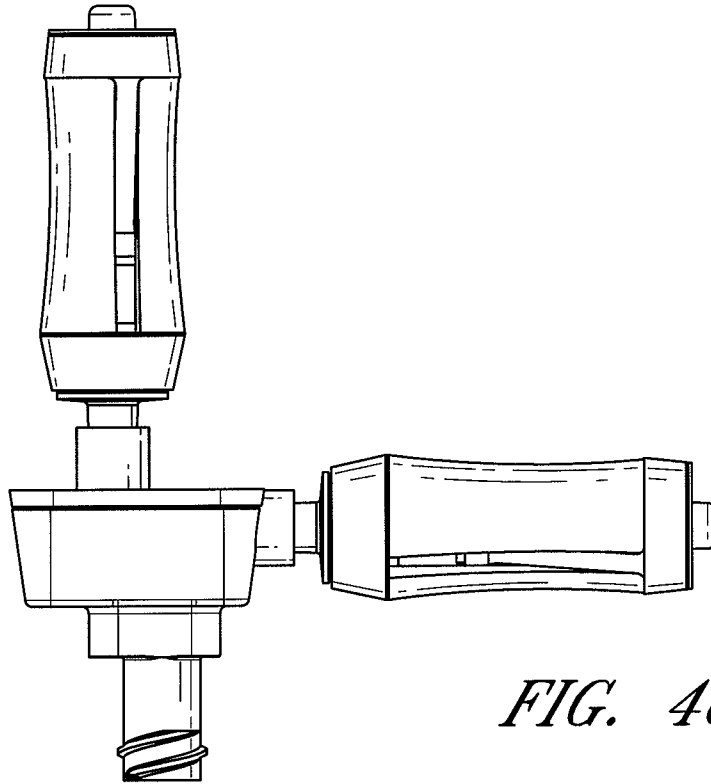


FIG. 44

*FIG. 45*



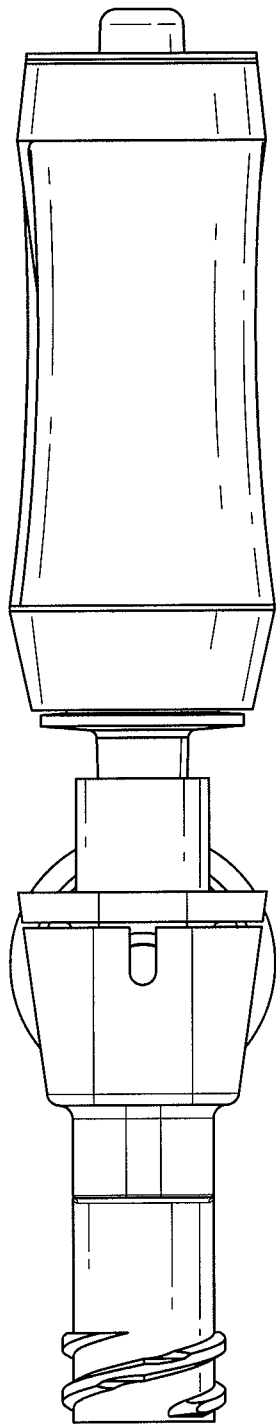


FIG. 47

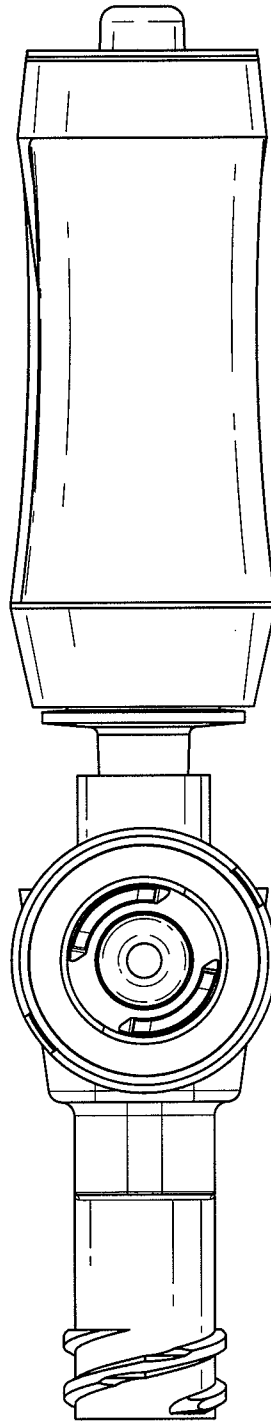


FIG. 48

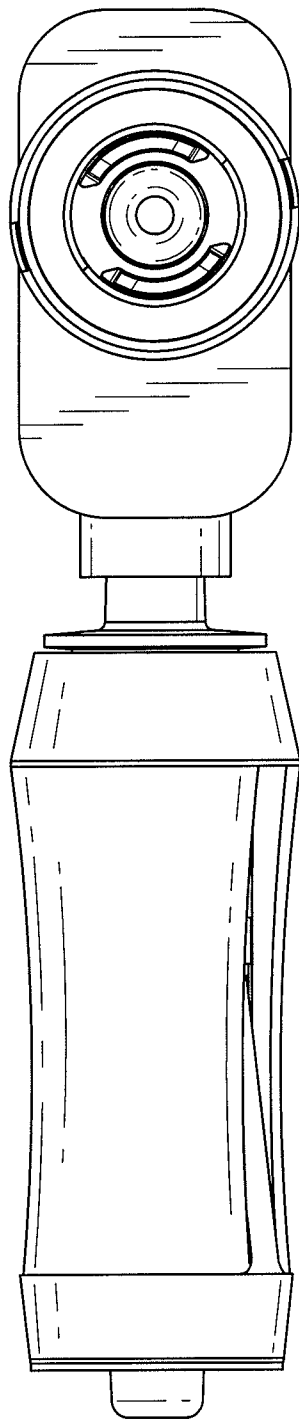


FIG. 49

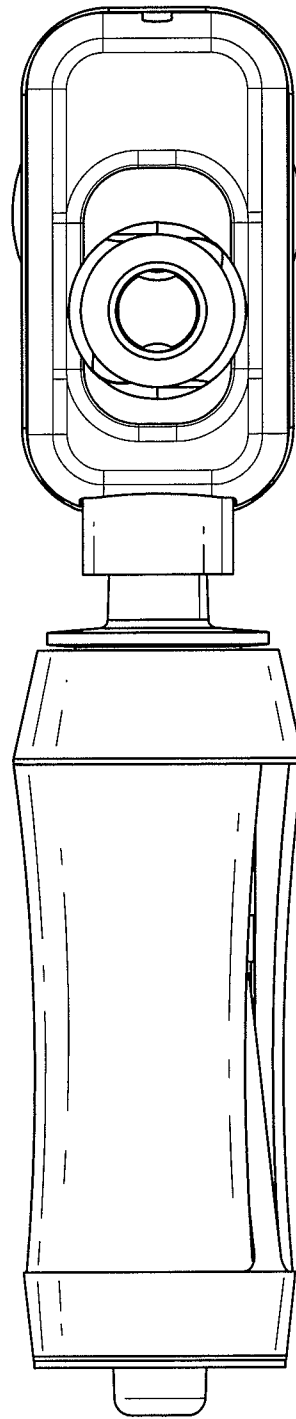


FIG. 50

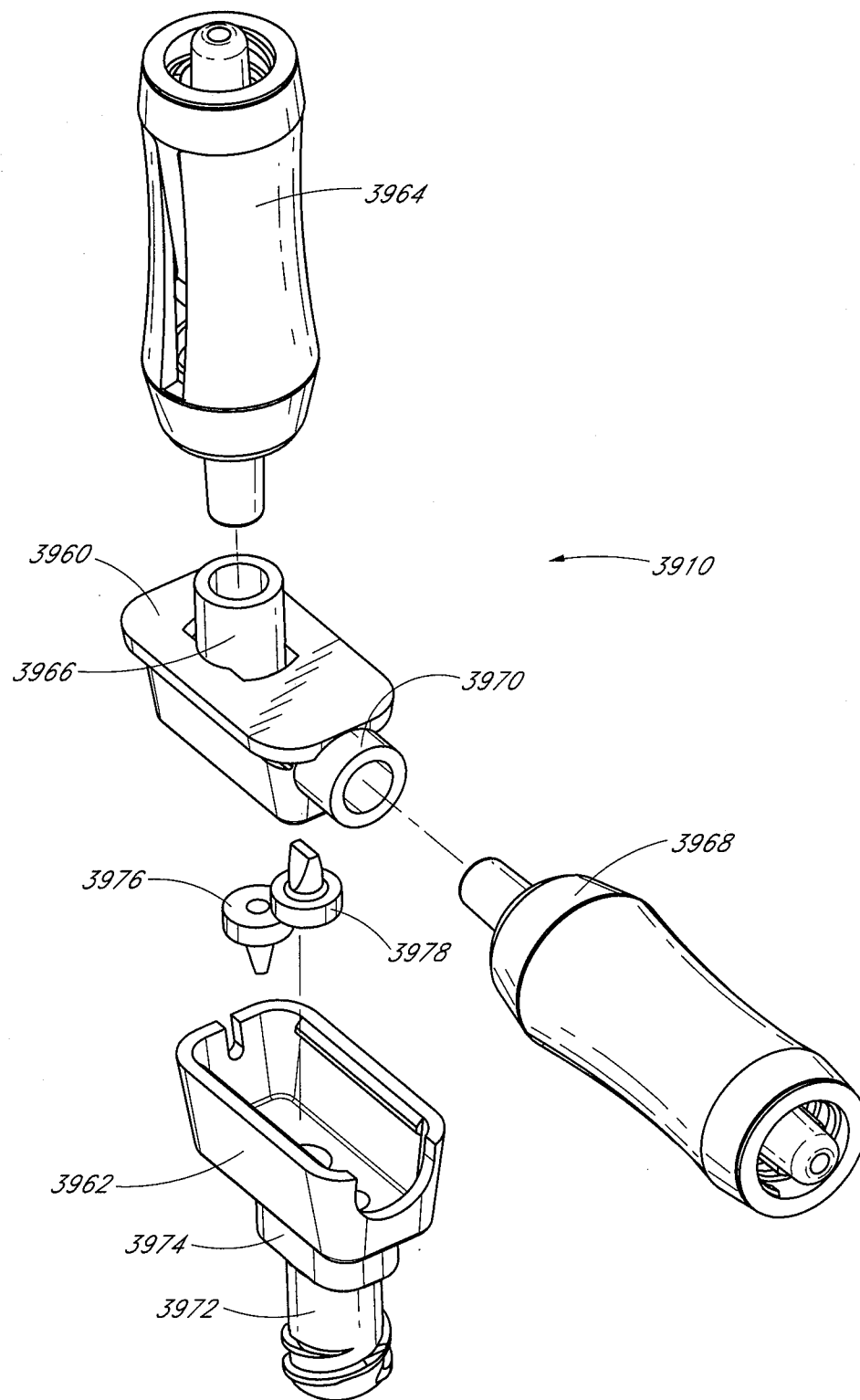


FIG. 52

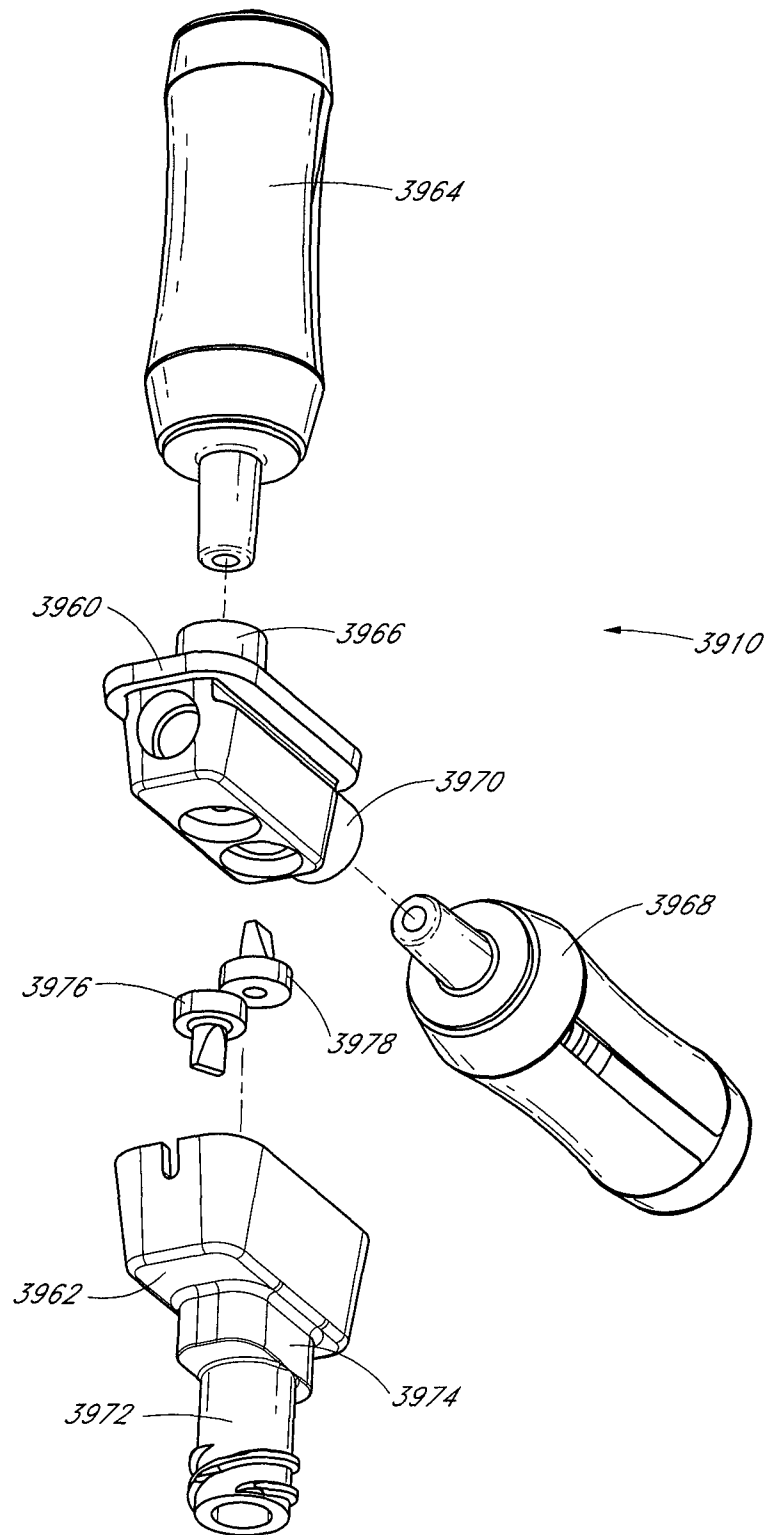


FIG. 53

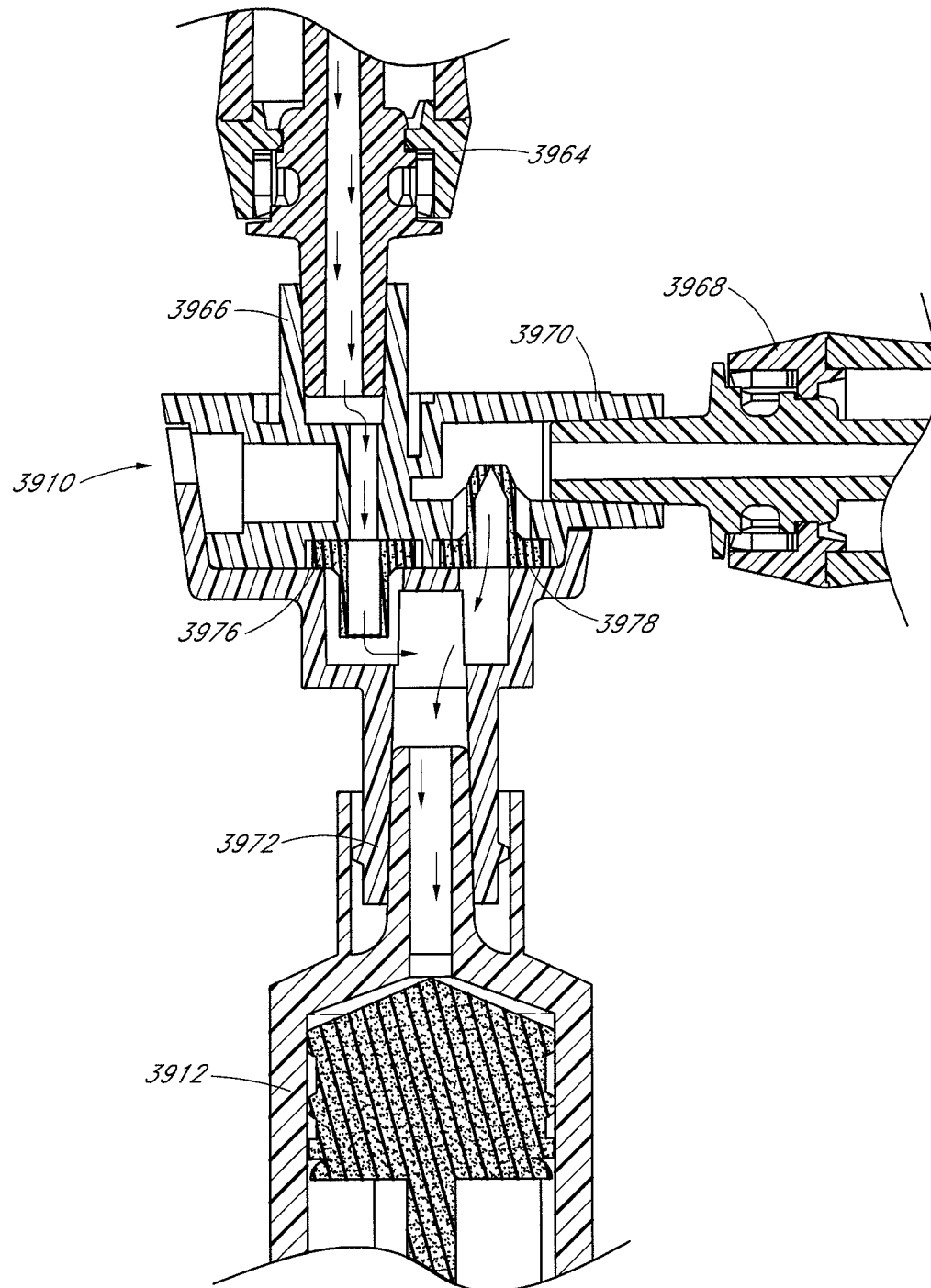
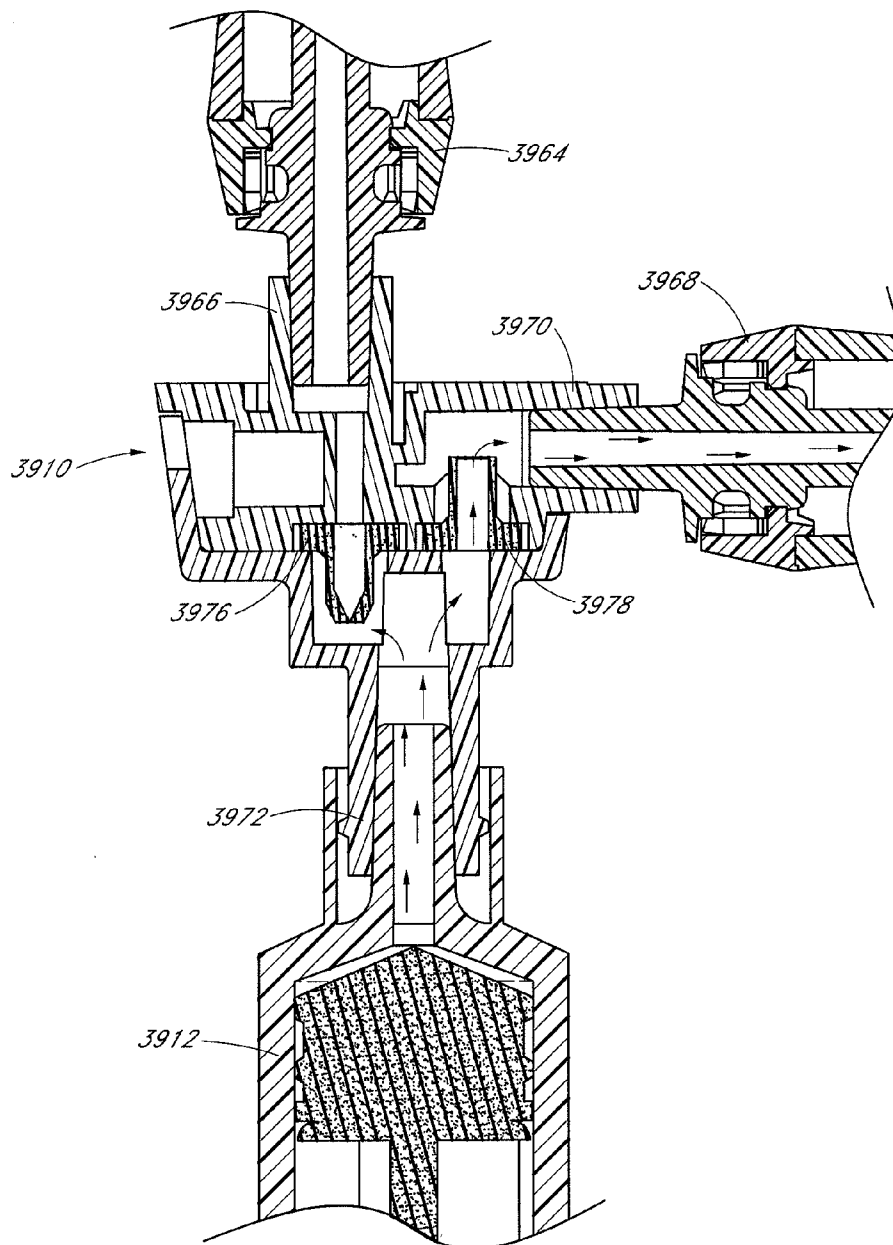


FIG. 54

*FIG. 55*

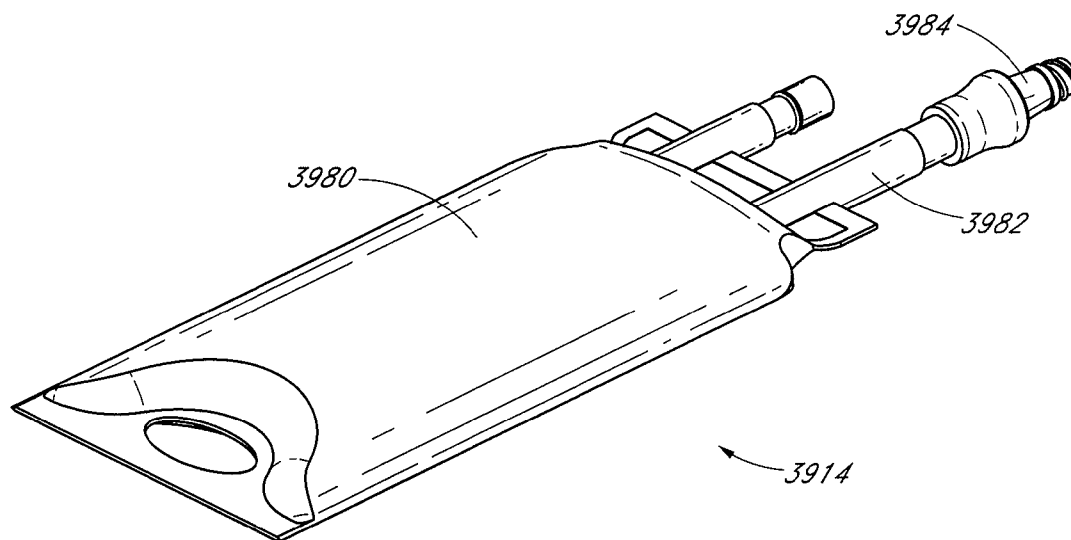


FIG. 56

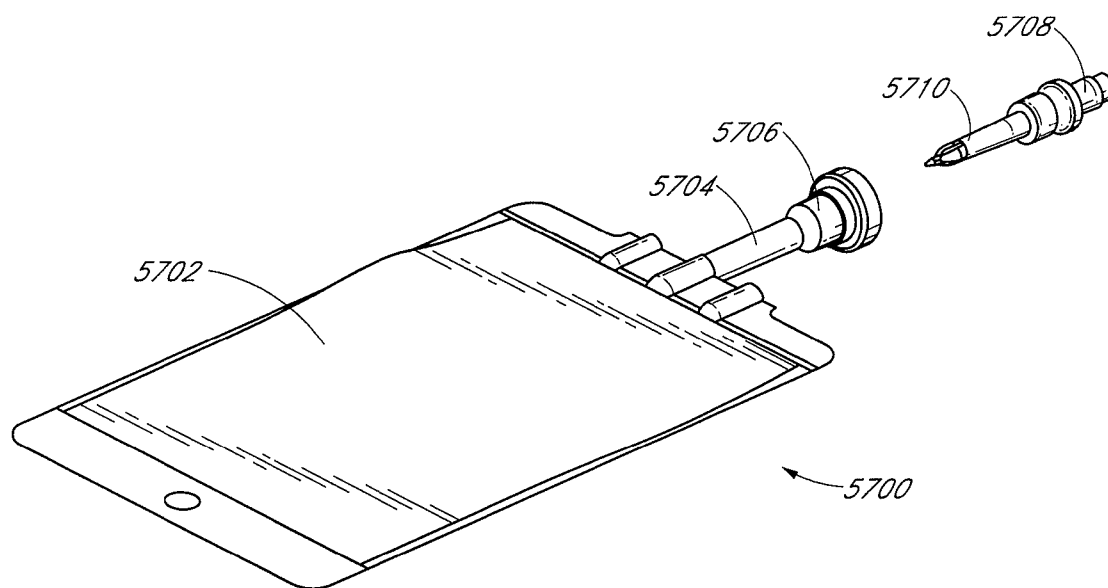


FIG. 57

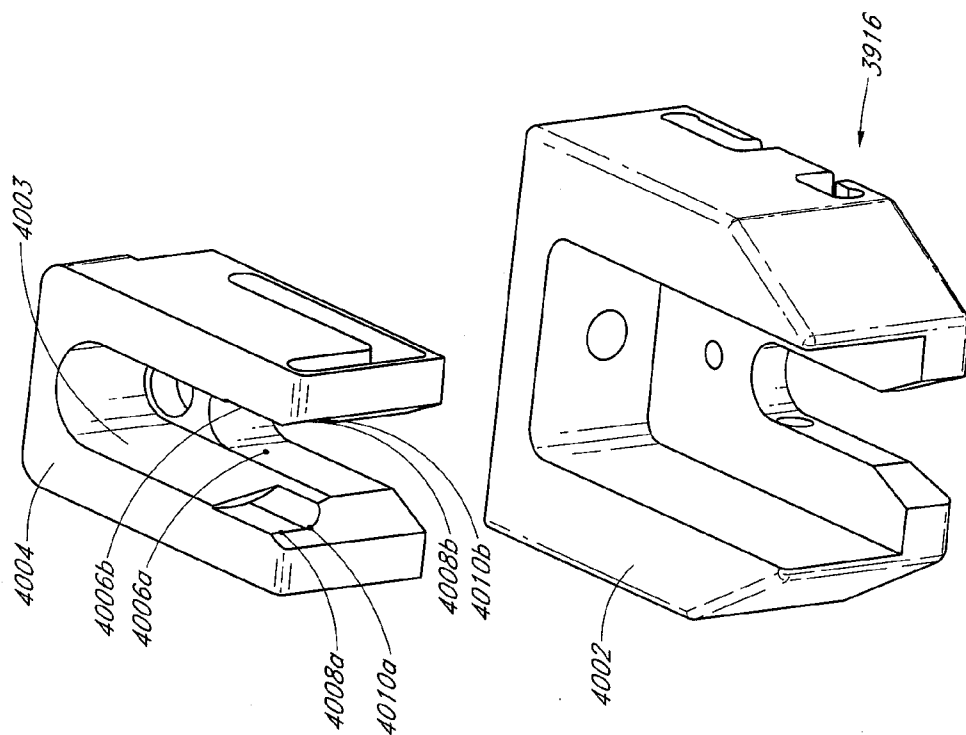


FIG. 59

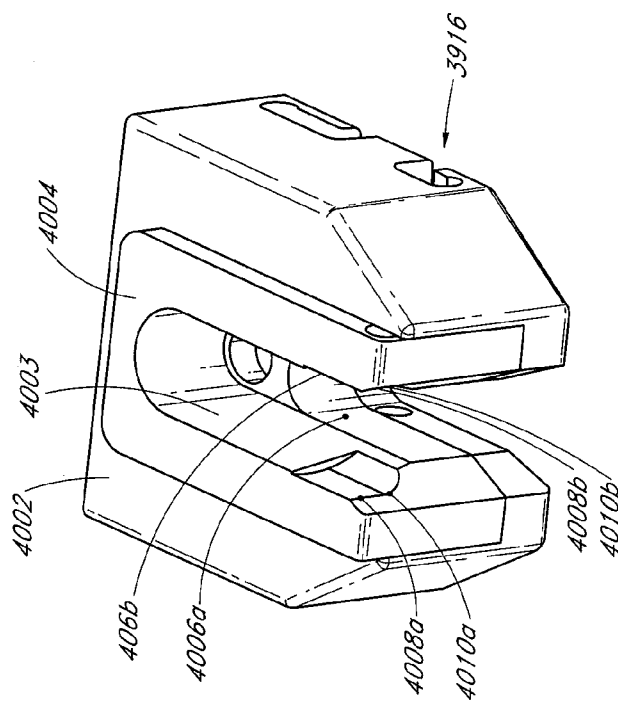


FIG. 58

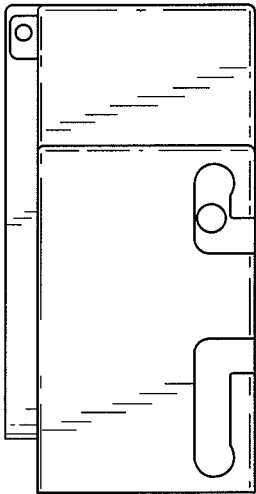


FIG. 60

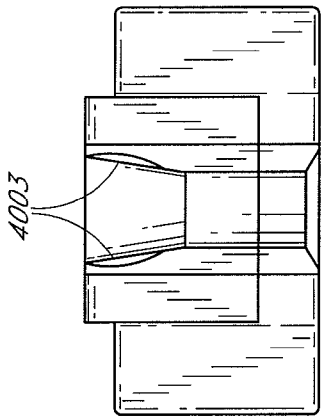


FIG. 61

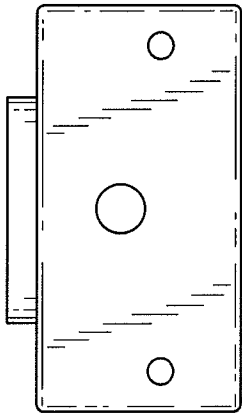


FIG. 62

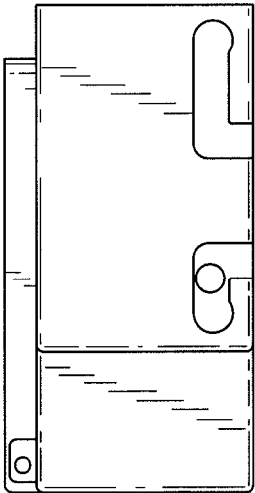


FIG. 63

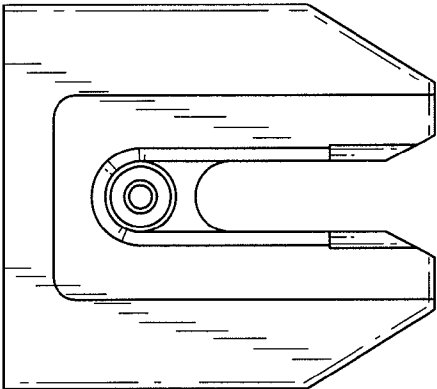


FIG. 64

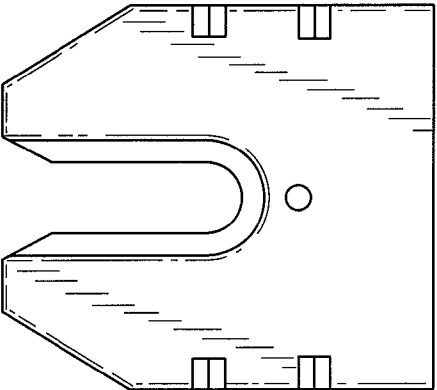


FIG. 65

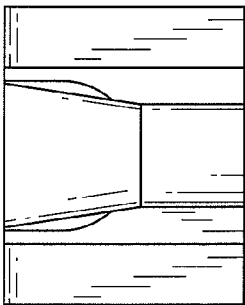


FIG. 66

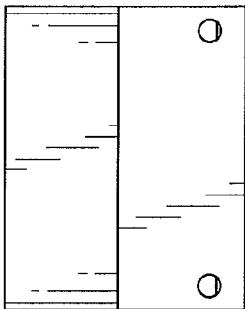


FIG. 67

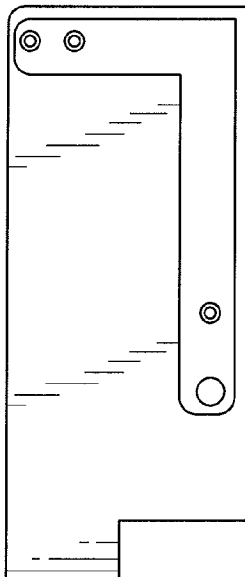


FIG. 68

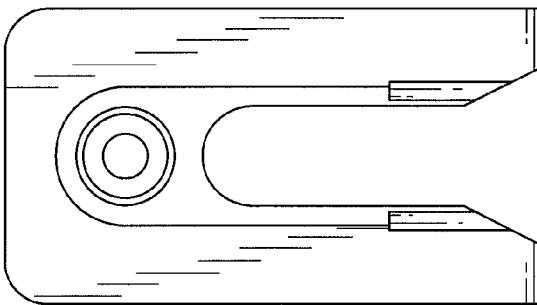


FIG. 69

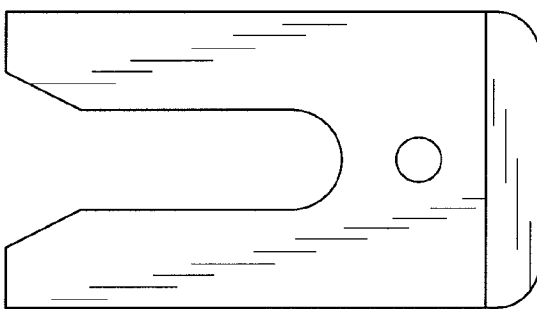


FIG. 70

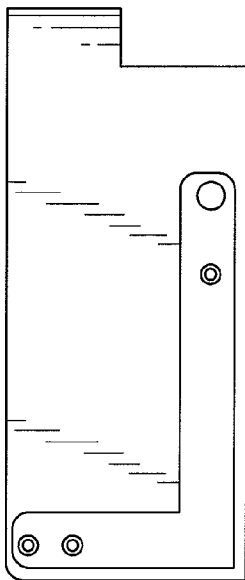


FIG. 71

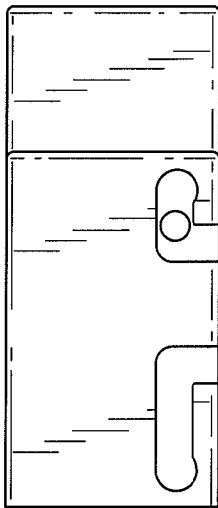


FIG. 72

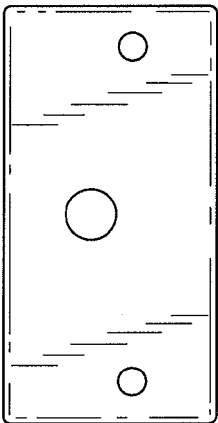


FIG. 73

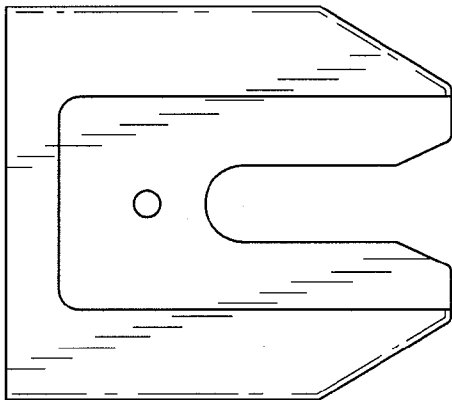


FIG. 74

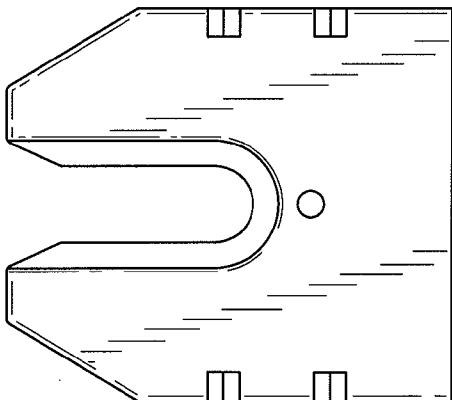


FIG. 75

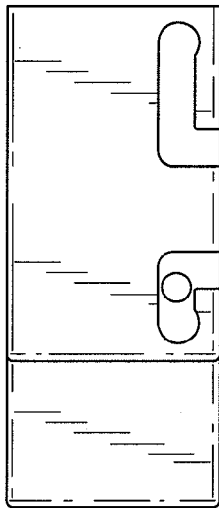


FIG. 76

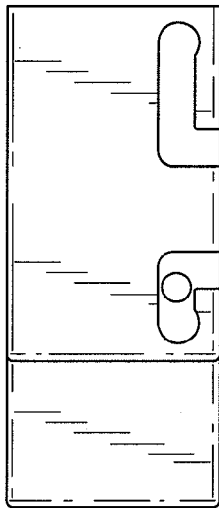


FIG. 77

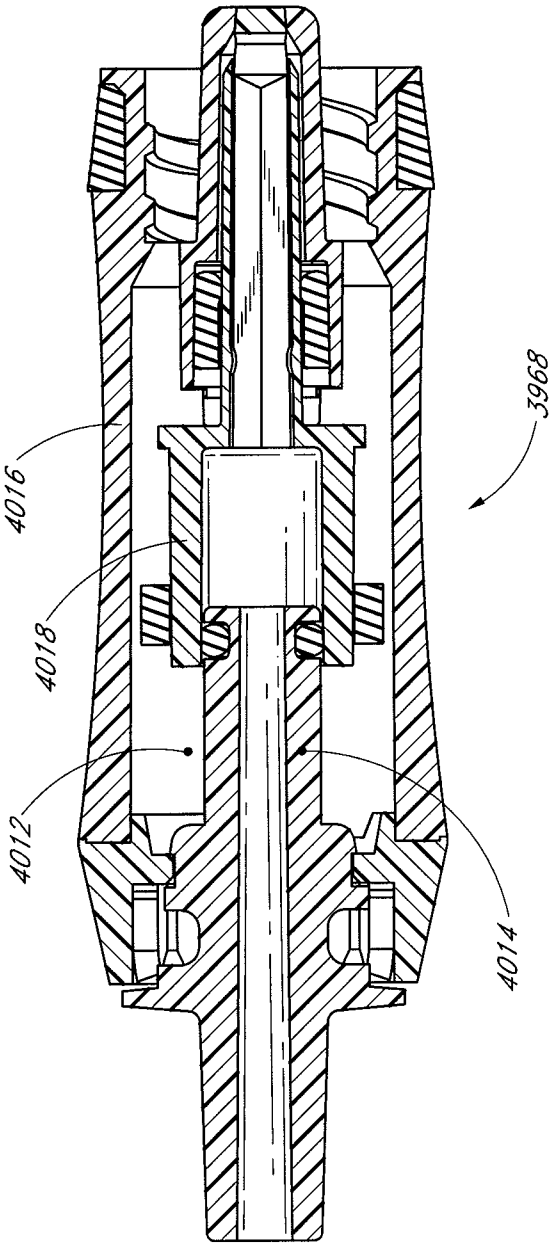


FIG. 78

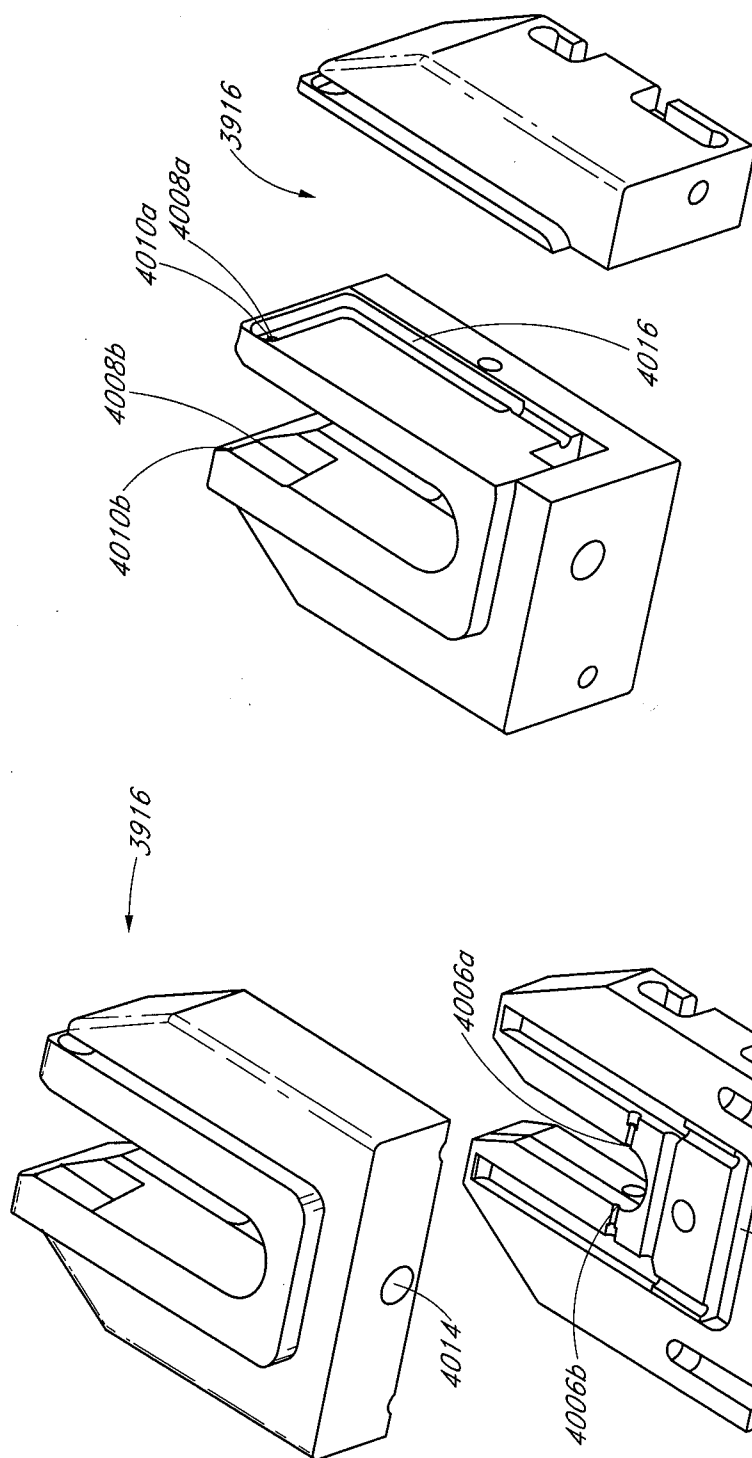


FIG. 80

FIG. 79

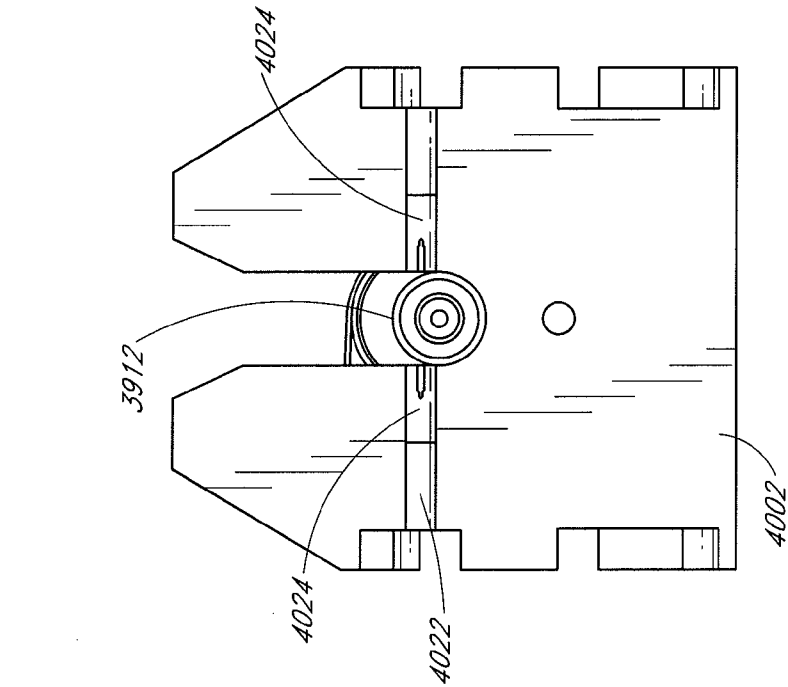


FIG. 81

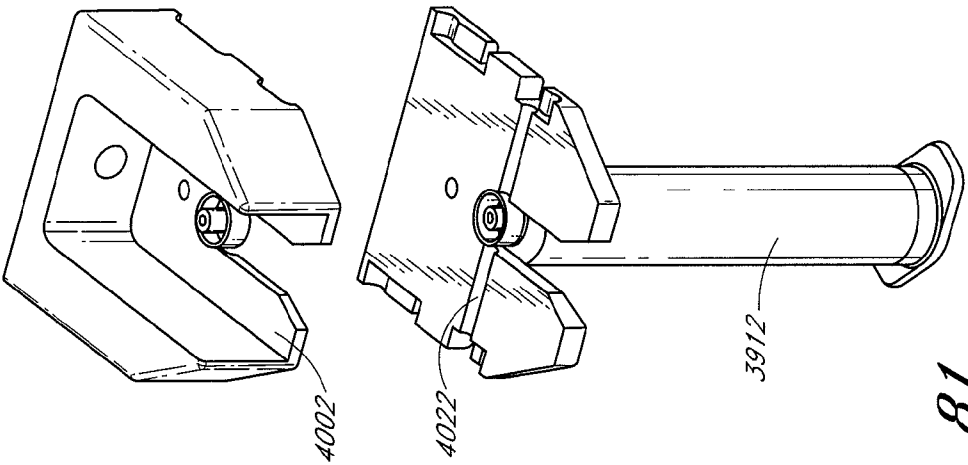


FIG. 82

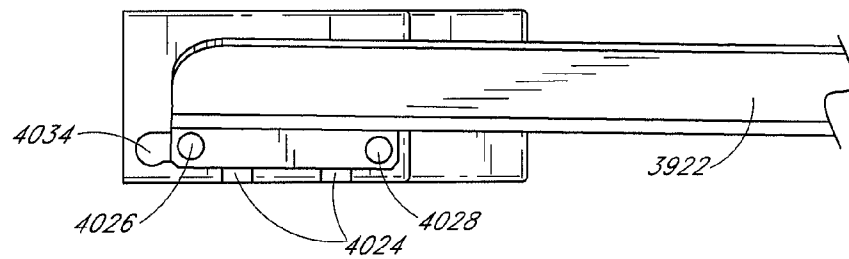


FIG. 83

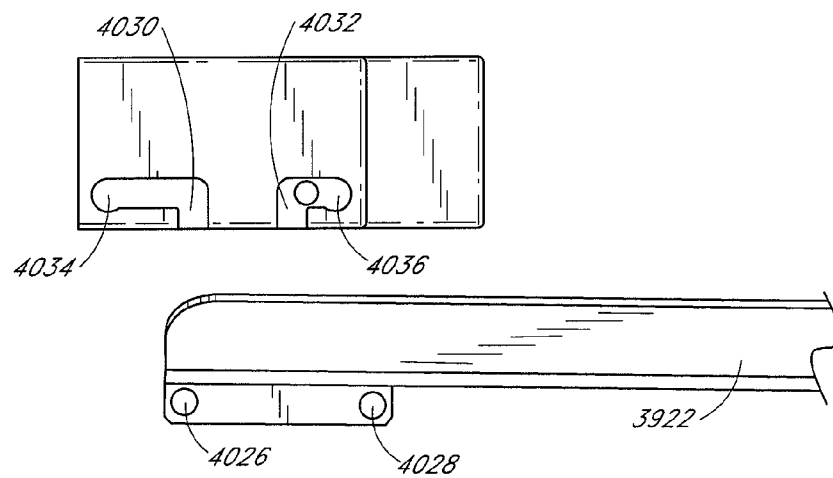
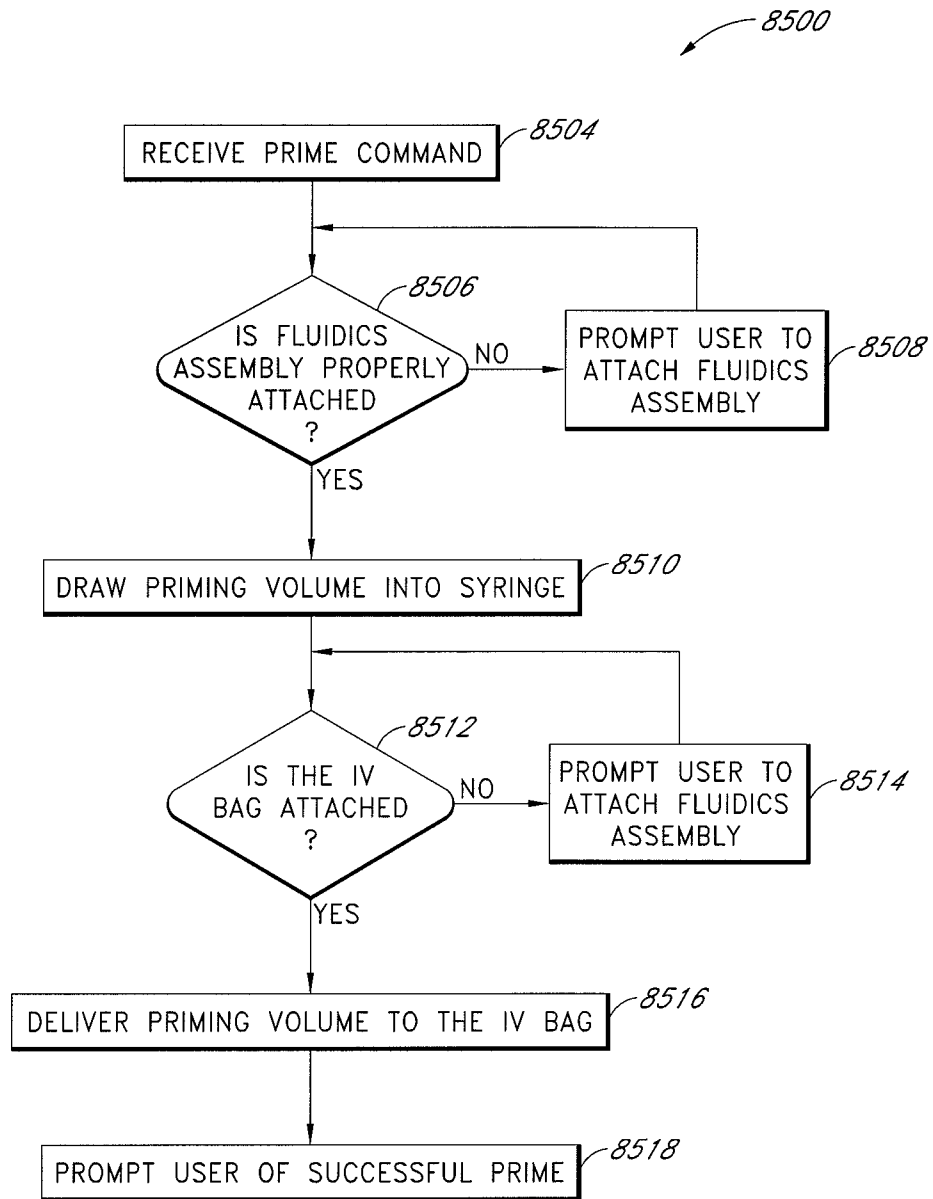
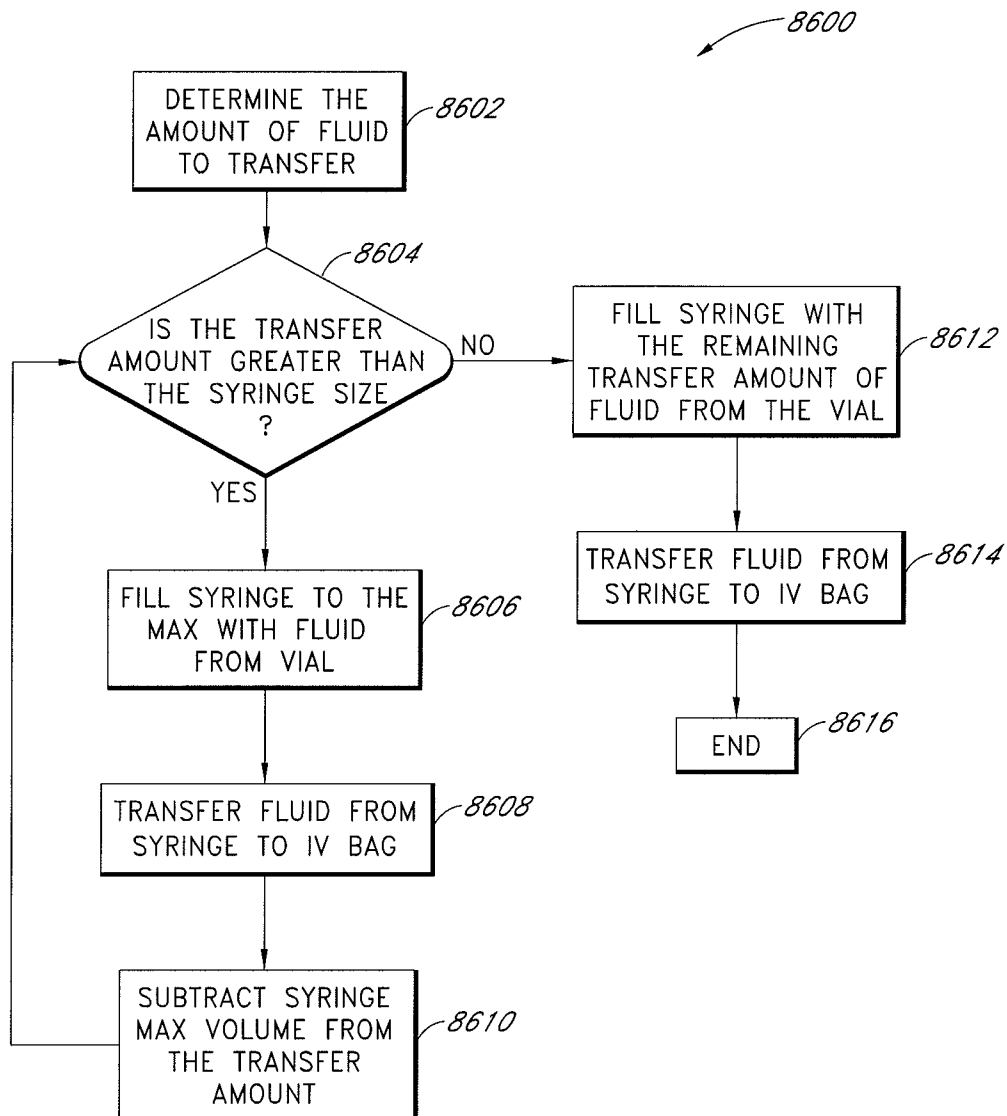
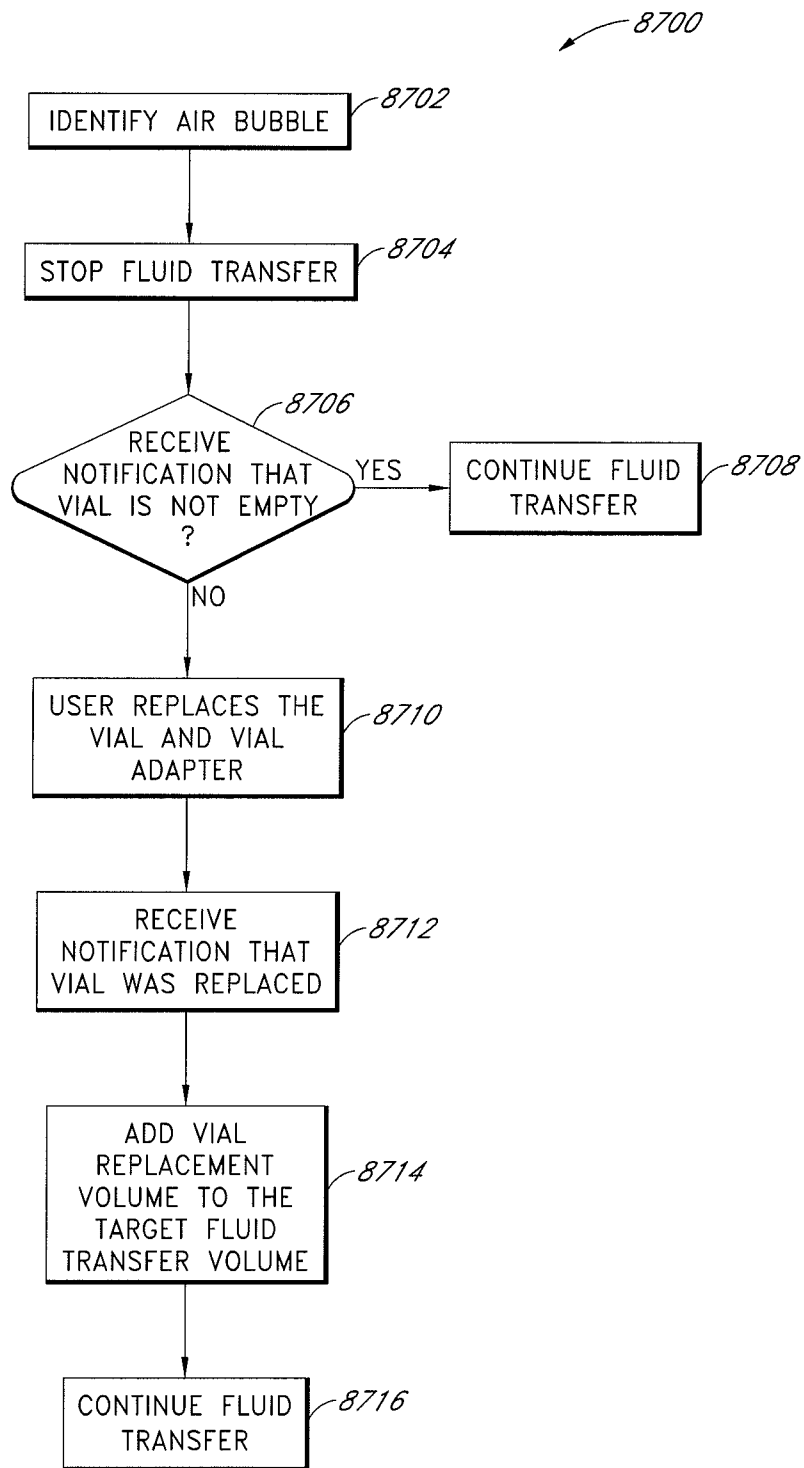
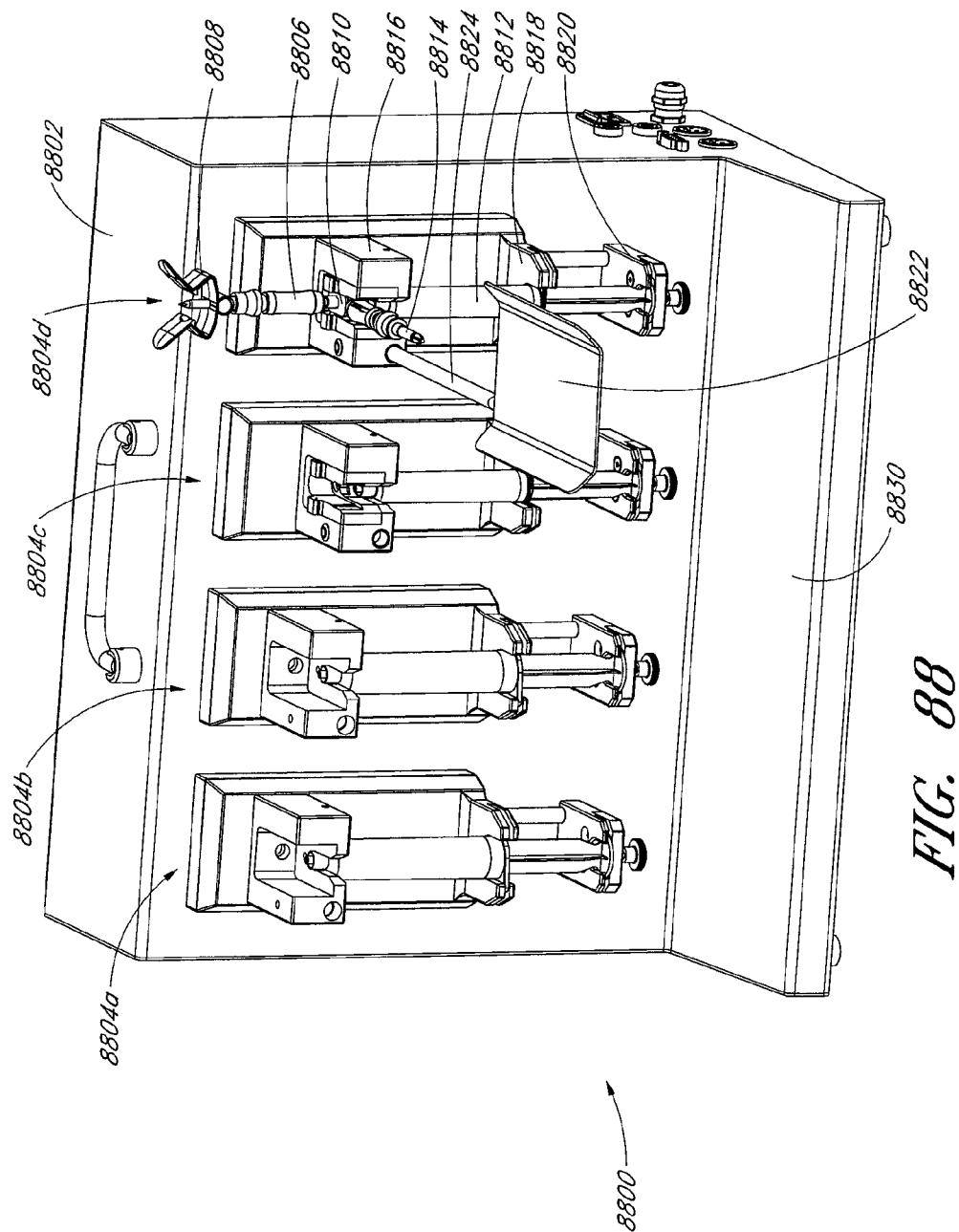


FIG. 84

*FIG. 85*

*FIG. 86*

*FIG. 87*



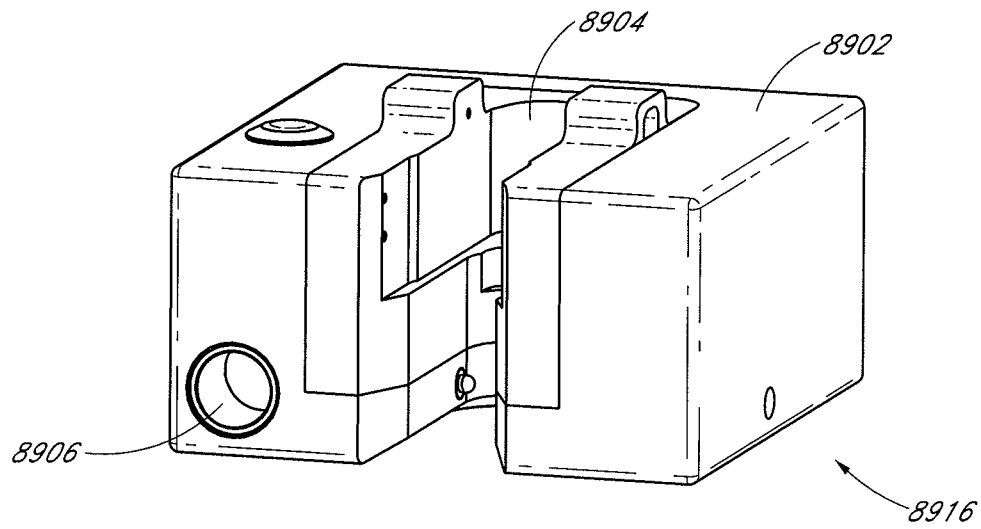


FIG. 89

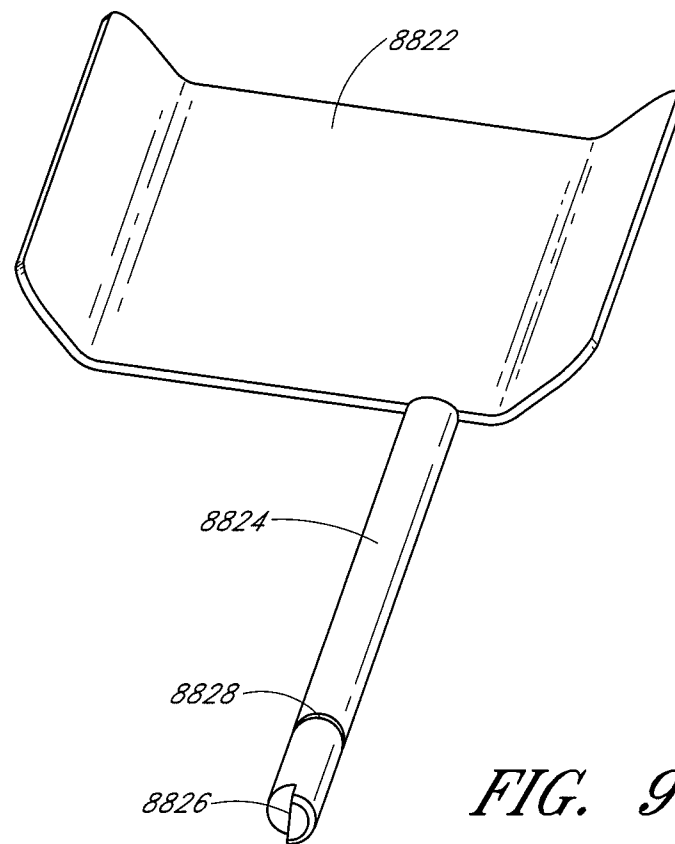
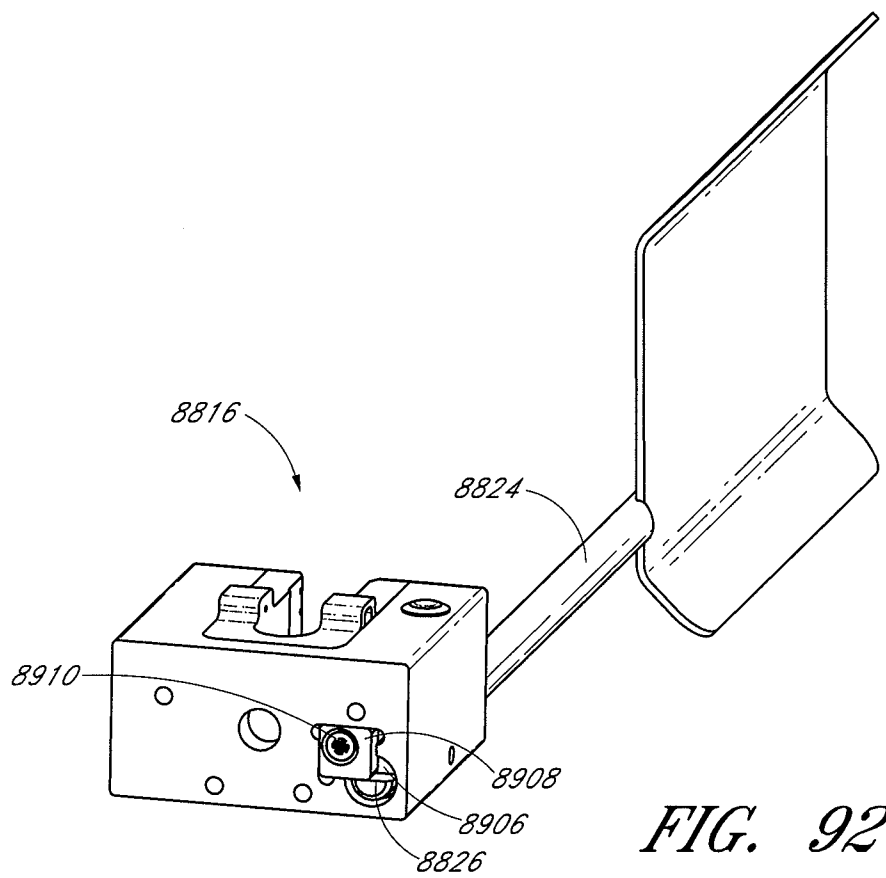
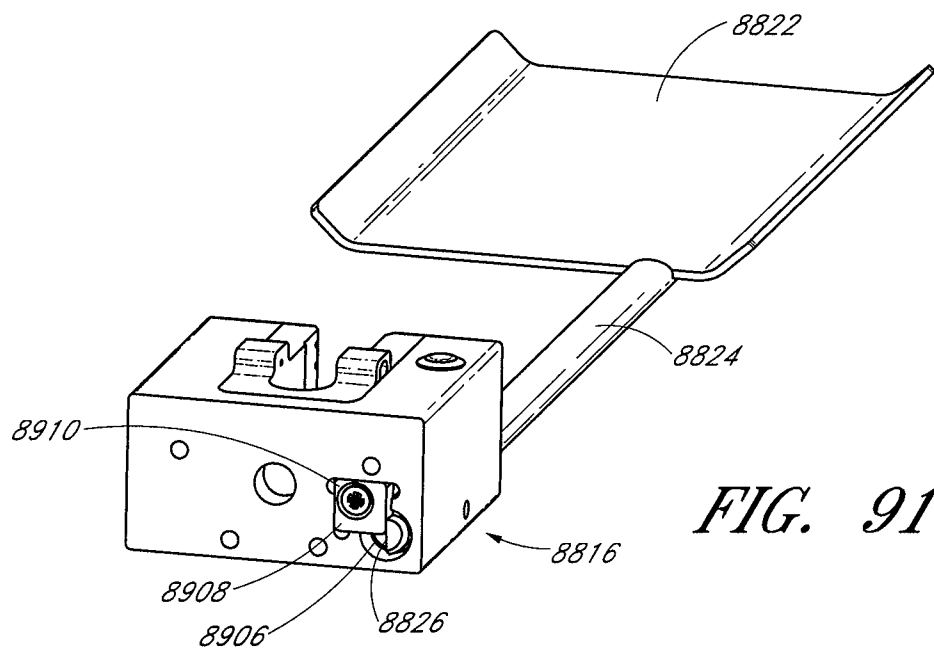


FIG. 90



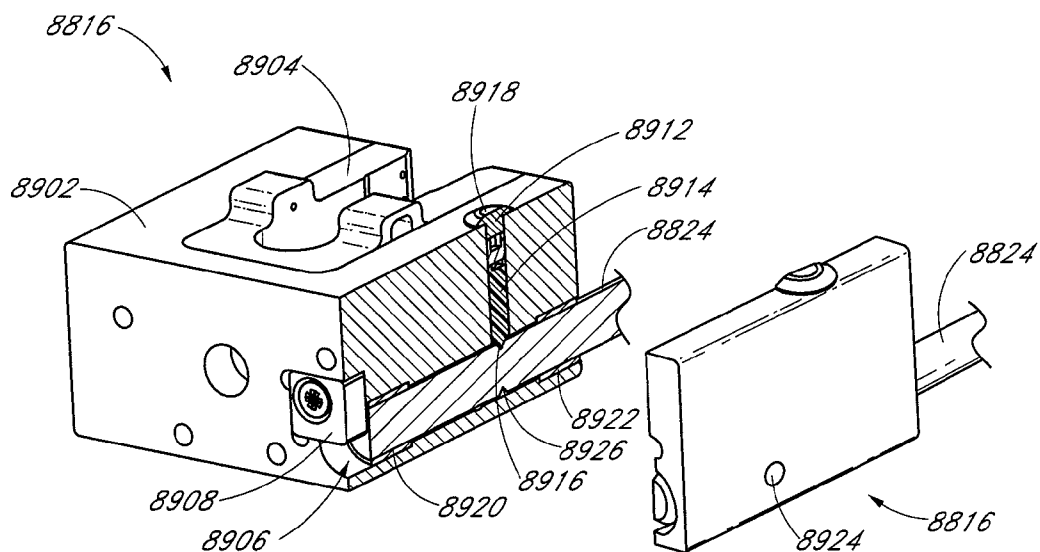


FIG. 93

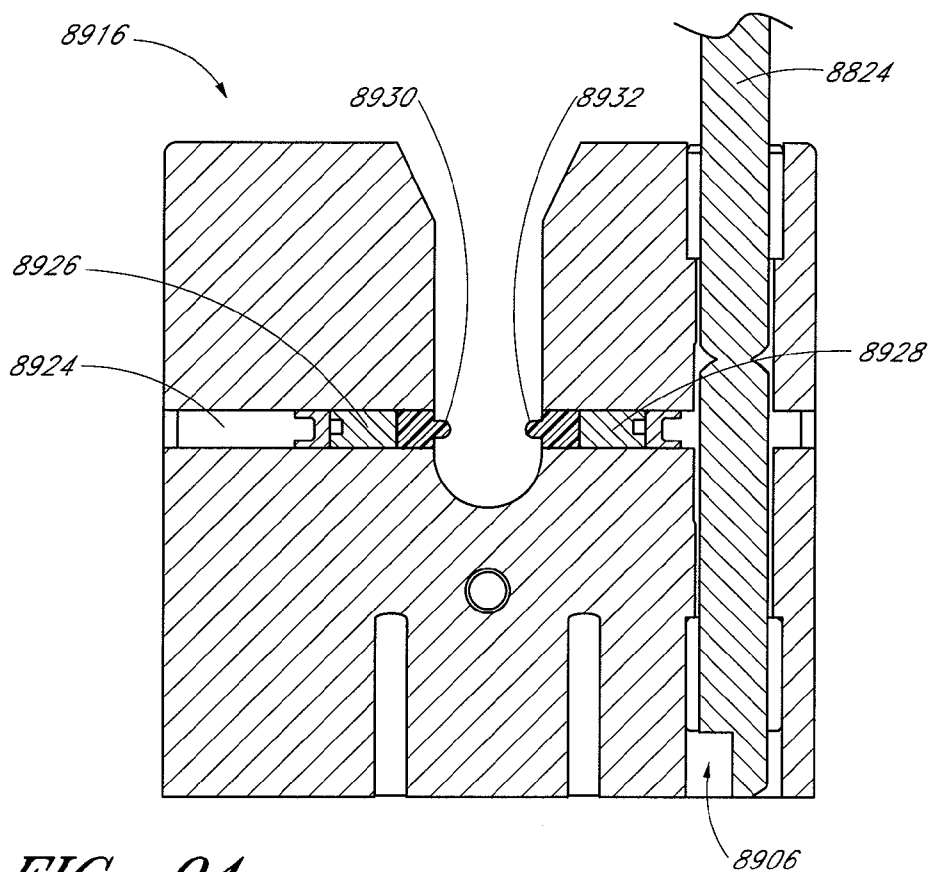


FIG. 94

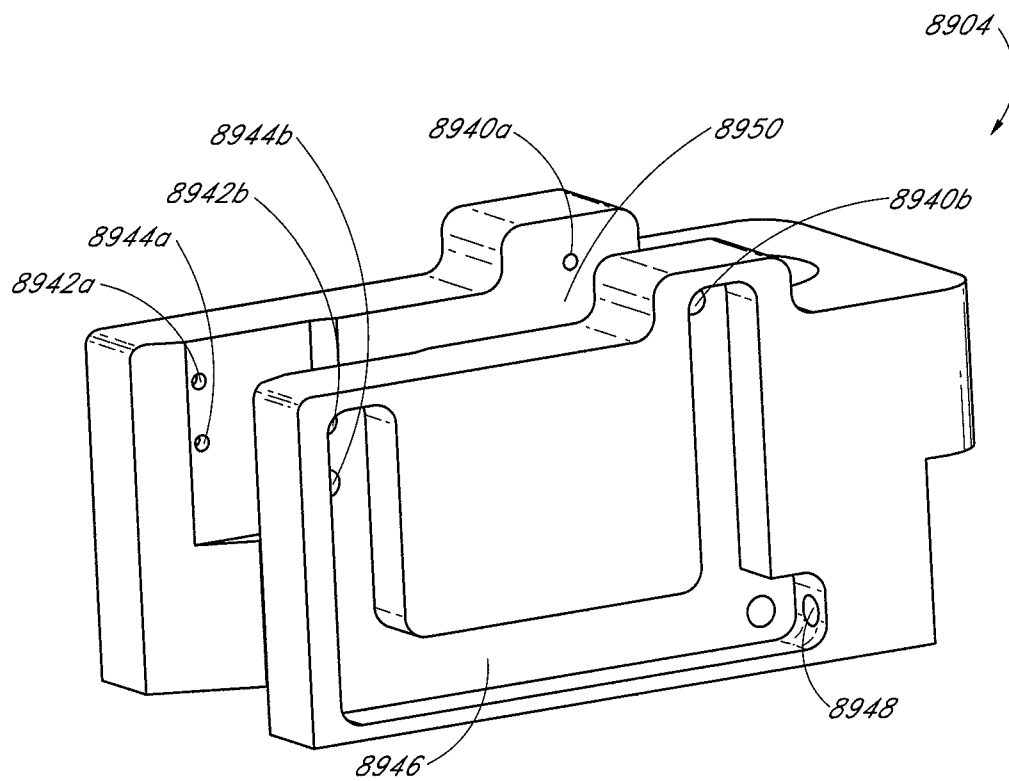


FIG. 95

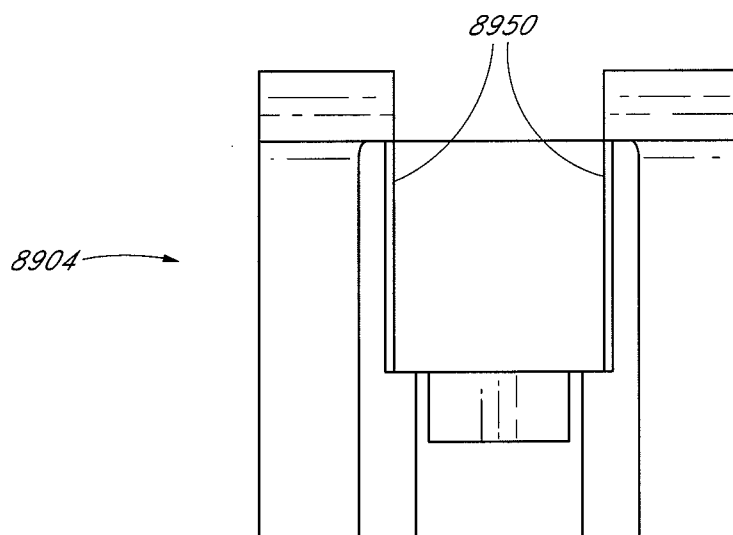


FIG. 96

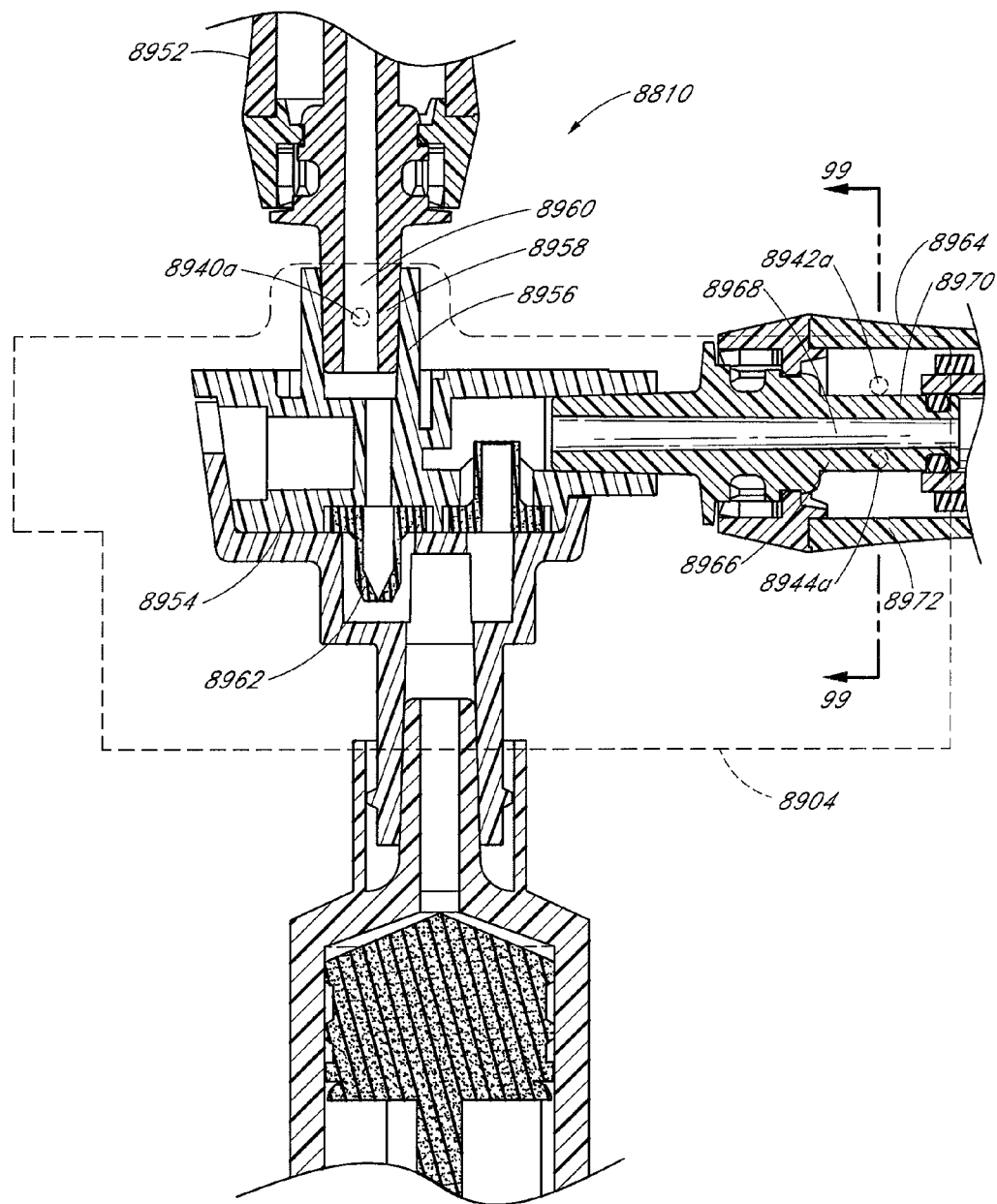
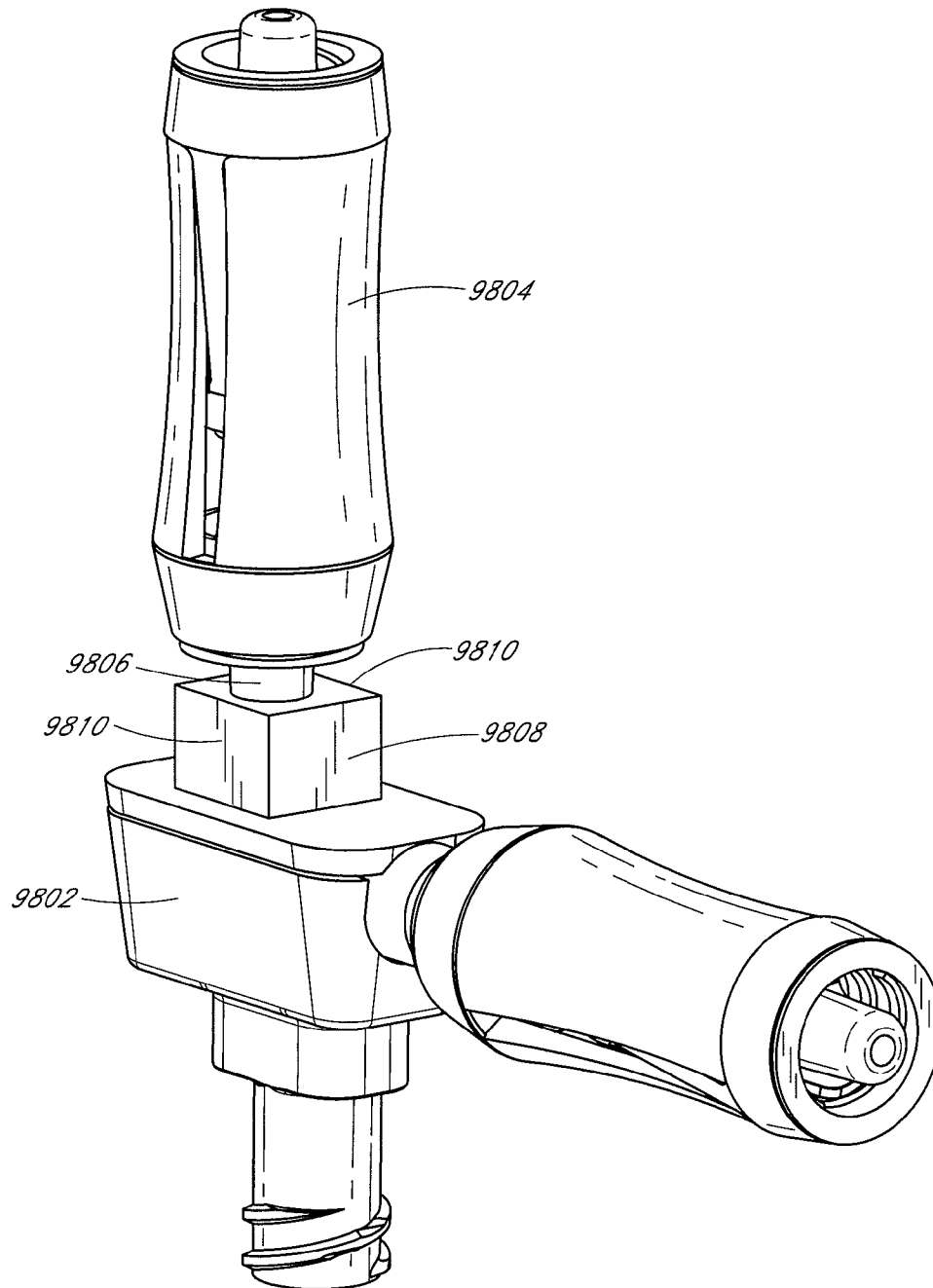


FIG. 97

*FIG. 98*

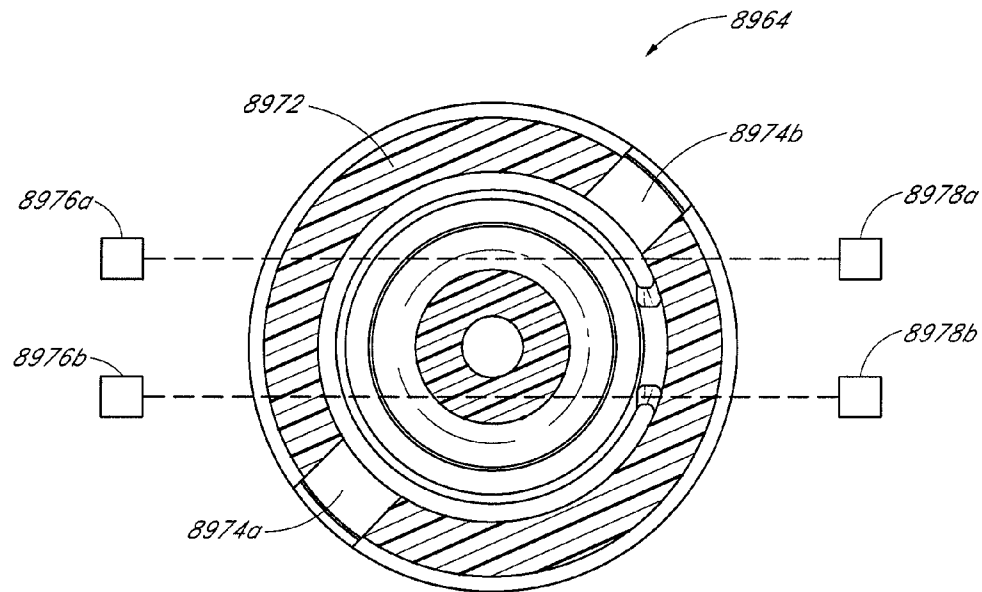


FIG. 99

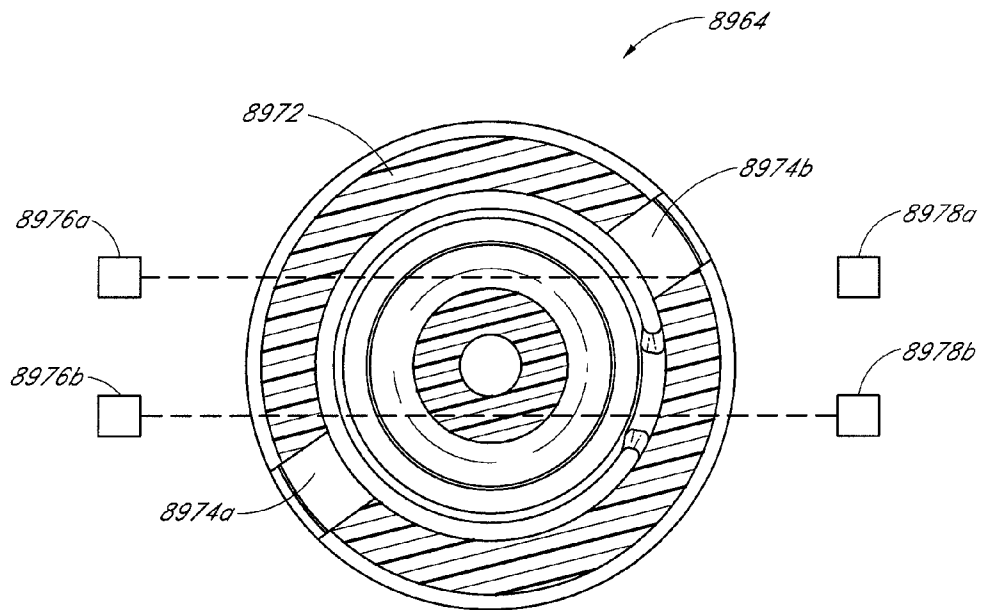


FIG. 100

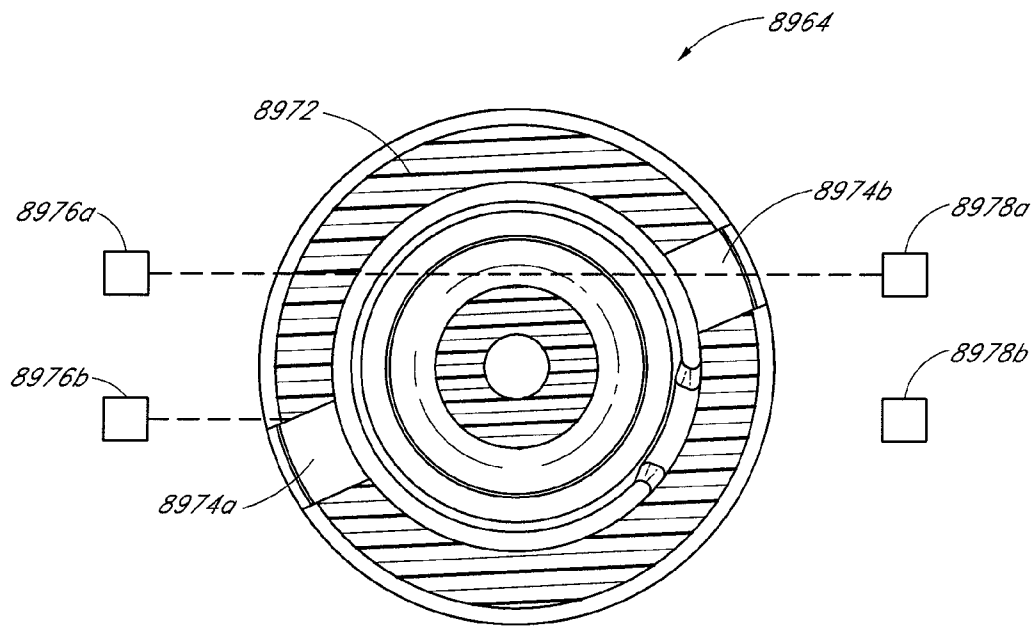


FIG. 101

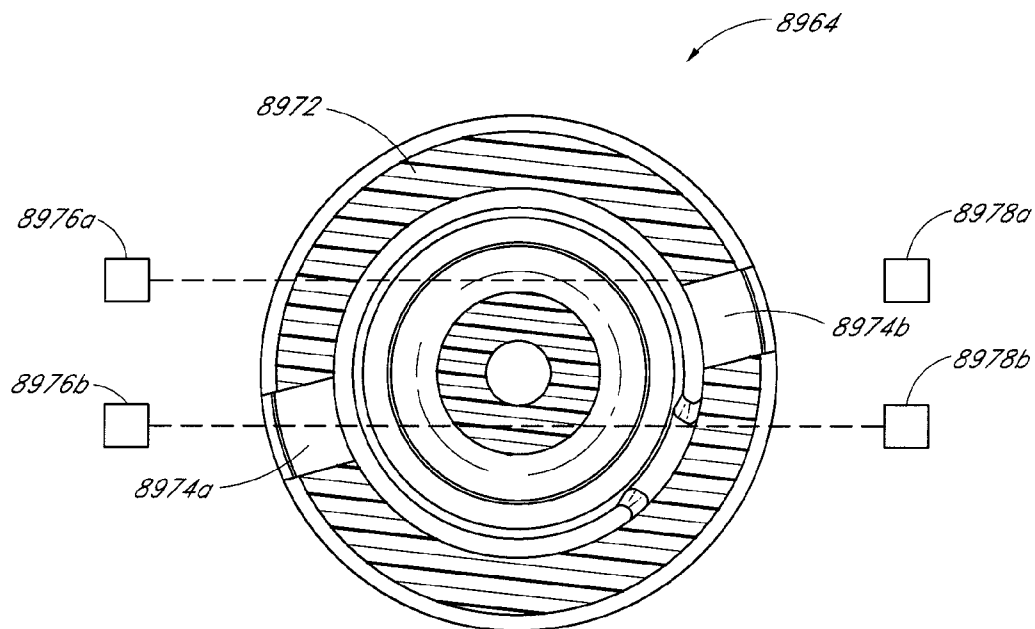


FIG. 102

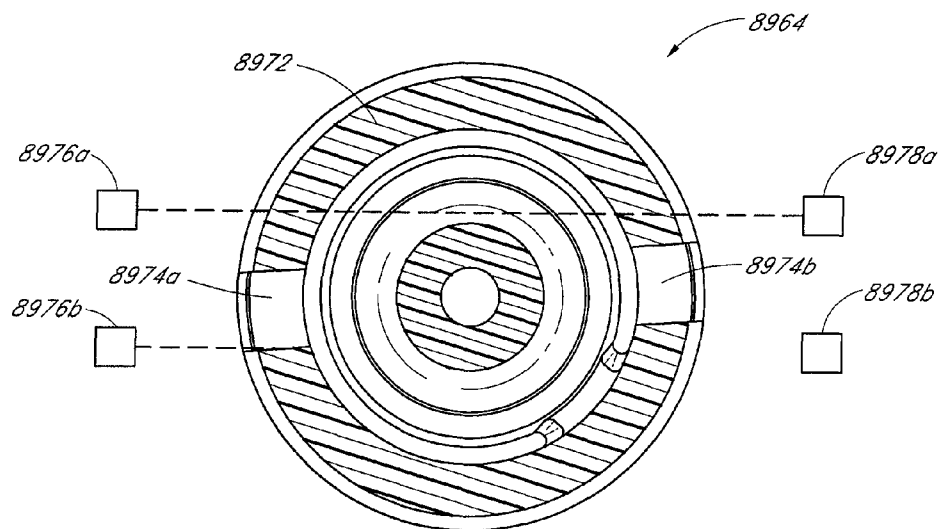


FIG. 103

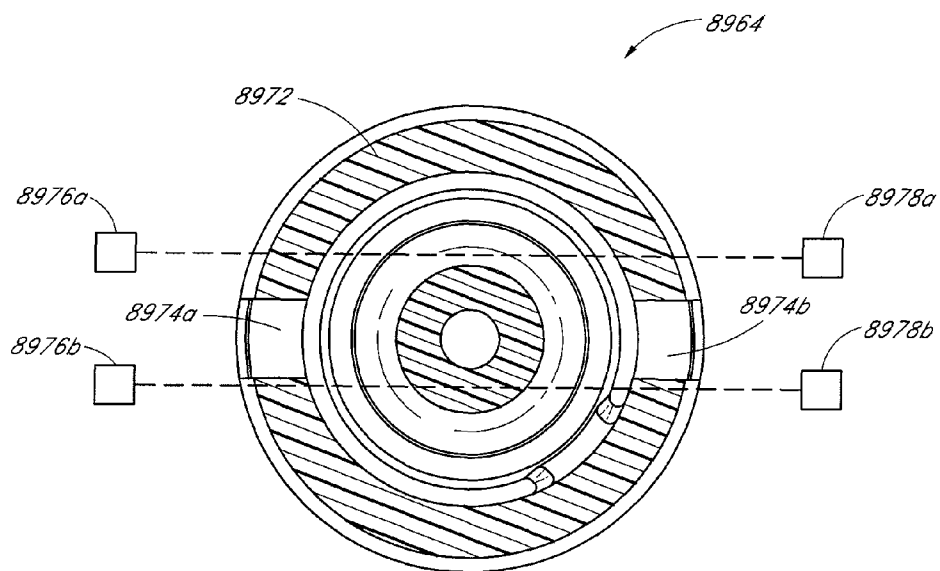
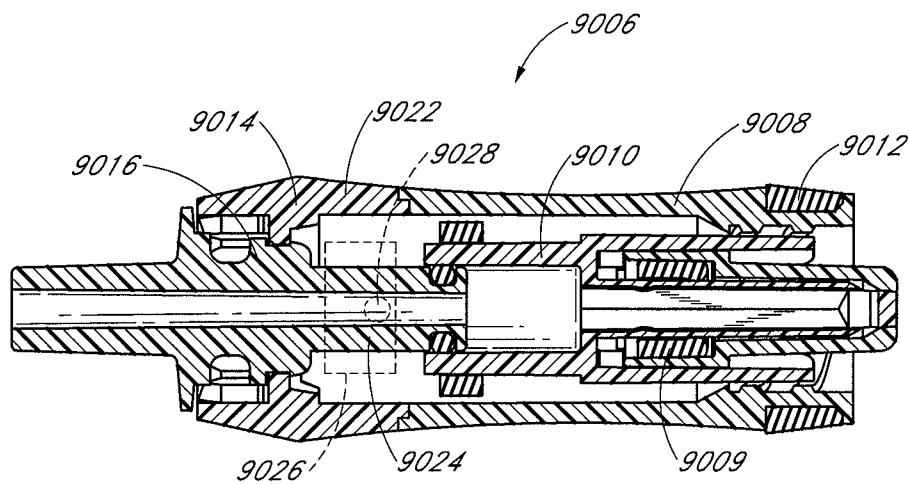
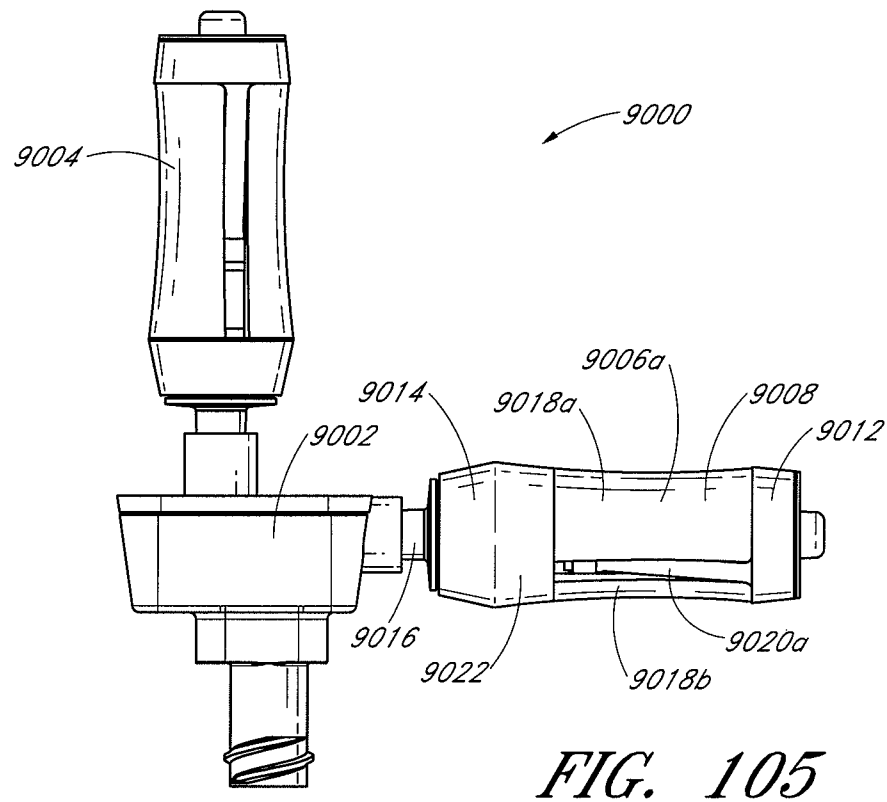


FIG. 104



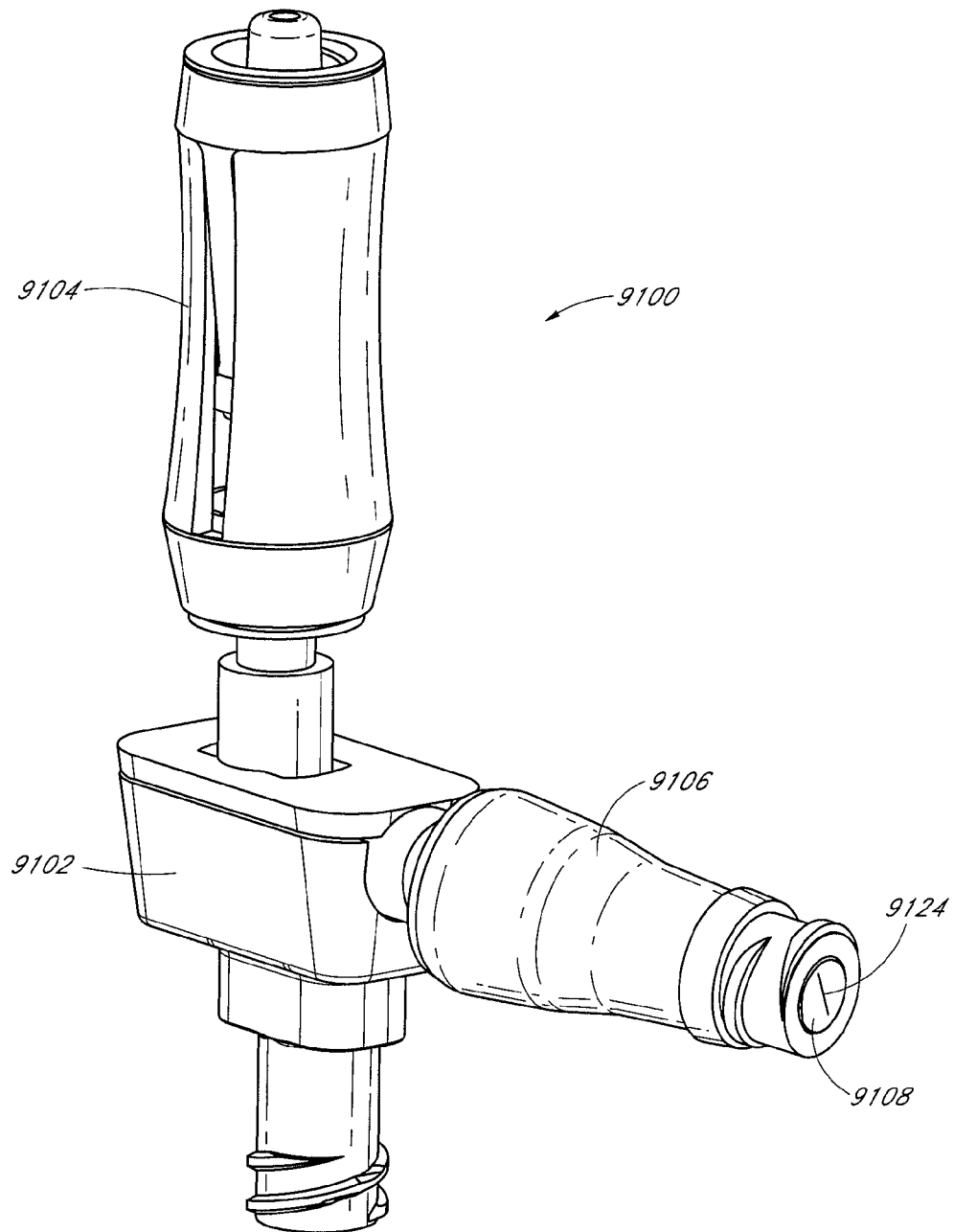


FIG. 107

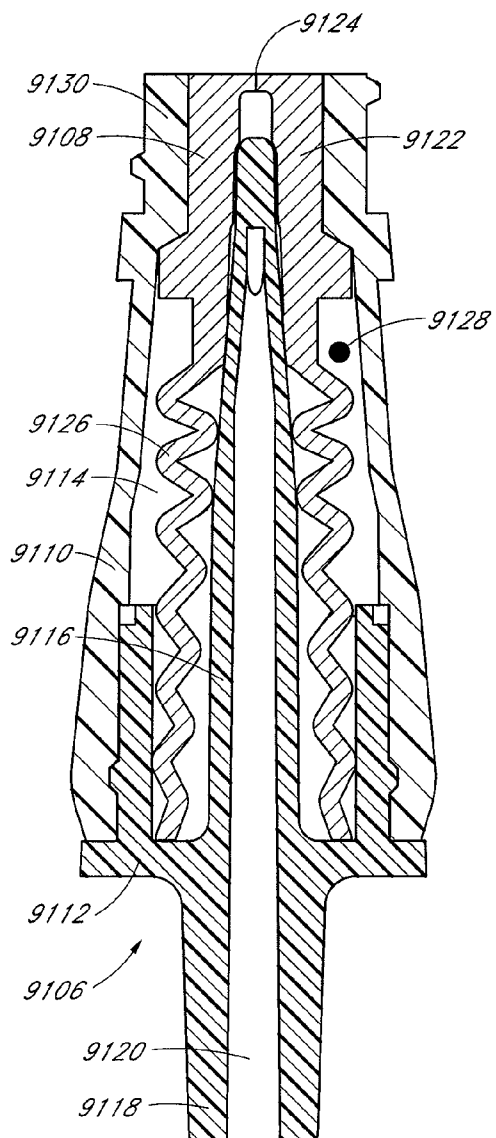


FIG. 108

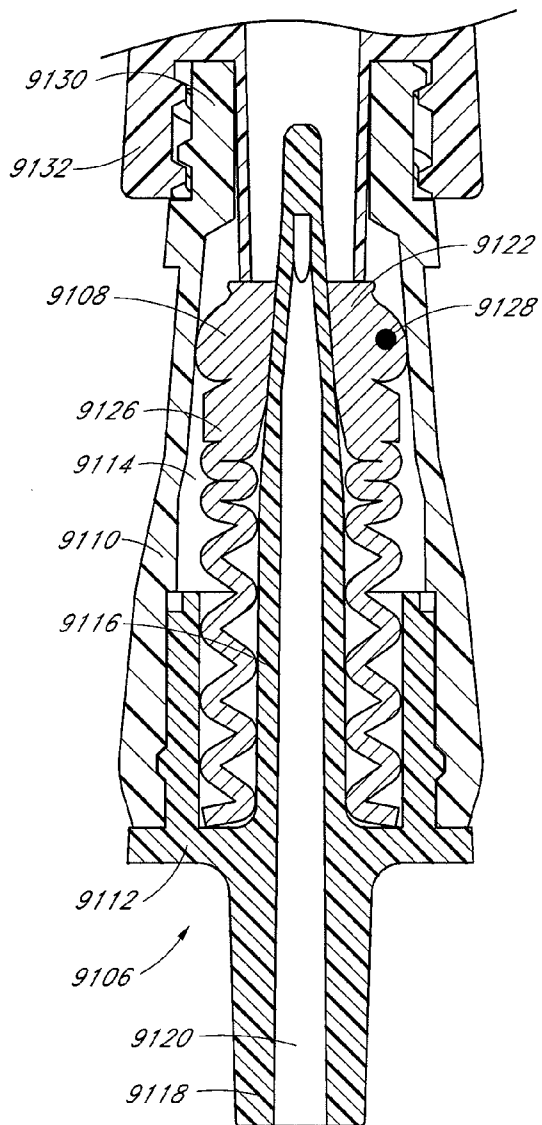


FIG. 109

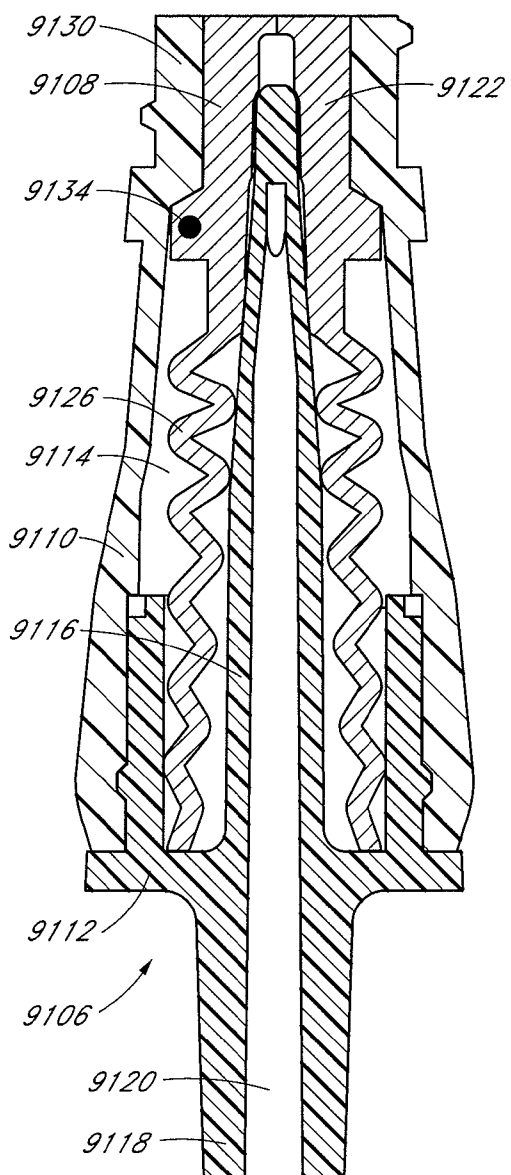


FIG. 110

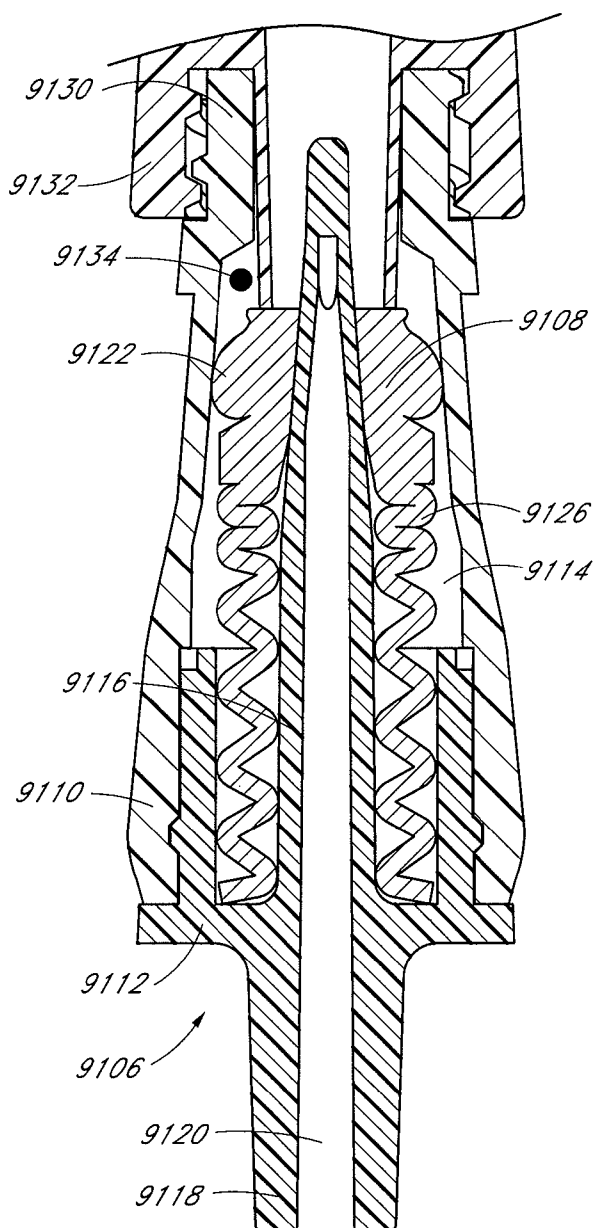


FIG. 111

1

FLUID TRANSFER DEVICES AND METHODS OF USE

RELATED APPLICATIONS

This application claims the benefit under 35 U.S.C. §119 (e) of U.S. Provisional Patent Application No. 61/229,701, filed Jul. 29, 2009, and entitled FLUID TRANSFER DEVICE, and U.S. Provisional Patent Application No. 61/354,648, filed Jun. 14, 2010, and entitled FLUID TRANSFER DEVICE, the entire contents of both of which are hereby incorporated by reference herein and made part of this specification for all that they disclose.

BACKGROUND OF THE DISCLOSURE

1. Field of the Disclosure

Some embodiments of the invention relate generally to devices and methods for transferring fluid and specifically to devices and method for transferring medical fluids.

2. Background of the Disclosure

In some circumstances it can be desirable to transfer one or more fluids between containers. In the medical field, it is often desirable to dispense fluids in precise amounts and to store and to transport potentially dangerous fluids. Current fluid transfer devices and methods in the medical field suffer from various drawbacks, including high cost, low efficiency, intensive labor demands, and excessive fluid or vapor leakage. Some embodiments disclosed herein overcome one or more of these disadvantages.

SUMMARY OF SOME EMBODIMENTS

Some embodiments disclosed herein related to devices for transferring precise amounts of fluid from a source container to a target container. In some embodiments, the fluid is first transferred from the source container through a connector to an intermediate measuring container (e.g., a syringe). The precisely measured amount of fluid can then be transferred from the intermediate measuring container to the target container.

In some embodiments, methods and devices for providing a substantially entirely closed system for the transfer of medical fluids between or among different medical fluid containers include a fluid transfer module that can be removably attached to an electronically controlled fluid dispensing system. The fluid transfer module can comprise first and second interfaces connected respectively to fluid source and fluid destination containers. The first and second interfaces can comprise selectively openable and closeable apertures that can substantially entirely prevent fluid within the fluid transfer module from escaping through the apertures when closed. An intermediate container can be part of or connected to the fluid transfer module. One or more valves within the fluid transfer module can permit fluid to move from the fluid source to the intermediate container, but can generally obstruct the fluid from moving from the intermediate container to the fluid source, and can permit fluid to move from the intermediate container to the fluid destination, but can generally obstruct the fluid from moving from the fluid destination to the intermediate container. In some embodiments, the fluid transfer module can be attached to an electronically controlled fluid dispensing system, and the fluid transfer module can include an interaction portion configured to permit the electronically controlled fluid dispensing system to indicate that at least a portion of the fluid transfer module is attached to the electronically controlled fluid dispensing system. In some

2

embodiments, the electronically controlled fluid dispensing system can include an interactive user interface and can be configured to dispense precise amounts of medical fluid.

BRIEF DESCRIPTION OF THE DRAWINGS

Certain embodiments of the invention will now be discussed in detail with reference to the following figures. These figures are provided for illustrative purposes only, and the embodiments are not limited to the subject matter illustrated in the figures.

FIG. 1 schematically shows an embodiment of an automated system for transferring precise amounts of fluid.

FIG. 2 schematically shows an embodiment of an automated system for compounding mixtures of precise amounts of fluid.

FIG. 3A is a perspective view of a subsystem for transferring fluid.

FIG. 3B is an exploded perspective view of the subsystem of FIG. 3A.

FIG. 4A is an exploded perspective view of the connector of FIG. 3A.

FIG. 4B is a cross sectional view of the connector of FIG. 4A.

FIG. 5A is a perspective view of the source connector portion of FIG. 4A adjacent to the vial of FIG. 3A.

FIG. 5B is another perspective view of the source connector portion of FIG. 4A and the vial of FIG. 3A.

FIG. 5C is a cross-sectional view of the source connector portion and vial of FIG. 5A in engagement.

FIG. 5D is a cross-sectional view of the source connector portion and vial of FIG. 5B in a subsequent stage.

FIG. 6A is a perspective view of the target connector portion of FIG. 4A.

FIG. 6B is an exploded perspective view of the target connector portion of FIG. 6A.

FIG. 6C is a top view of a housing portion of the target connector portion.

FIG. 6D is a cross-sectional view of the target connector portion and the female connector in an unengaged configuration.

FIG. 6E is a cross-sectional detail view of the target connector portion and the female connector in an engaged configuration.

FIG. 7A is a perspective view of the syringe connector portion of FIG. 4A adjacent to the syringe of FIG. 3A.

FIG. 7B is a top view of the syringe connector portion and the syringe of FIG. 7A in engagement.

FIG. 7C is a cross-sectional view of the syringe connector portion and syringe of FIG. 7A in engagement.

FIG. 8A is a perspective view of the source check valve of FIG. 4B.

FIG. 8B is another perspective view of the source check valve of FIG. 8A.

FIG. 9A is an exploded cross sectional view of the source connector portion and main body of FIG. 4A and the source check valve of FIG. 8A.

FIG. 9B is a cross sectional view of the source connector portion, main body, and source check valve shown in FIG. 9A in an assembled configuration.

FIG. 10A is a side view of the main body coupled to the source connector portion of FIG. 4A.

FIG. 10B is a cross sectional view of the source connector portion of FIG. 4A and the source check valve of FIG. 8A disposed therein.

FIG. 10C is a partial cross-sectional view of the source connector and source check valve shown in FIG. 10B.

3

FIG. 10D is a side cross sectional view showing the source connector portion and the source check valve of FIG. 10B.

FIG. 11 is a side cross sectional view of the source check valve of FIG. 10B positioned against a side wall of a chamber.

FIG. 12 is another side cross sectional view of the source check valve of FIG. 10B positioned against a side wall of a chamber.

FIG. 13A is an exploded cross sectional view of the main body, target connector portion, and target check valve of FIG. 4B.

FIG. 13B is a cross sectional view of the main body, target connector portion, and target check valve of FIG. 13A.

FIG. 14A is a cross sectional view of the fluid transfer system of FIG. 3A with the source check valve in an open configuration and the target check valve in a closed configuration.

FIG. 14B is a cross sectional view of the fluid transfer system of FIG. 3A with the source check valve in a closed configuration and the target check valve in an open configuration.

FIG. 15 is a perspective view of an automated system for transferring fluid having multiple transfer stations.

FIG. 16A is a perspective view of a transfer station of the system shown in FIG. 15.

FIG. 16B is a side view of the fluid transfer system shown in FIG. 15.

FIG. 16C is a front view of the transfer station shown in FIG. 16A.

FIG. 17 is a perspective view of the top connector piece of the transfer station shown in FIG. 16A with the top portion thereof removed to show a light source and photodetector disposed therein.

FIG. 18 is a cross sectional view of the syringe and connector of FIG. 15 showing regions where the light from the light source of FIG. 17 can intersect the connector.

FIG. 19A is a perspective view of another embodiment of a top connector piece.

FIG. 19B is an exploded perspective view of the top connector piece of FIG. 19A.

FIG. 19C is a side view of a connector for use in transferring fluid.

FIG. 19D is a cross sectional view of the connector of FIG. 19C in which the target connector portion is closed.

FIG. 19E is a cross sectional view of the connector of FIG. 19C in which the target connector portion is open.

FIG. 20 is a perspective view schematically showing another embodiment of an automated fluid transfer system wherein the system includes a support bar assembly attached to the housing.

FIG. 21 is a side view of an attachment piece and arm of FIG. 20.

FIG. 22 is a partial perspective view schematically showing another embodiment of an automated fluid transfer system wherein one or more of the transfer stations include a support arm.

FIG. 22A is a perspective view of a fluid transfer system that includes a support tray for supporting an IV bag.

FIG. 23 is a flowchart that shows an embodiment of a method of operation for an automated fluid transfer system.

FIG. 24 is a flowchart that shows an embodiment of a method for transferring fluid.

FIG. 25 is a flowchart that shows an embodiment of a method for confirming the successful transfer of fluid by checking the IV bag weight.

FIG. 26 is a cross sectional view of another embodiment of a connector for transferring fluid.

4

FIG. 27A is a perspective view of another embodiment of a connector for transferring fluid.

FIG. 27B is another perspective view of the connector of FIG. 27A.

FIG. 28A is an exploded perspective view of the connector of FIG. 27A.

FIG. 28B is another exploded perspective view of the connector of FIG. 27A.

FIG. 29A is a perspective view of a duckbill check valve.

FIG. 29B is another perspective view of the duckbill check valve of FIG. 29A.

FIG. 29C is a cross sectional view of the duckbill check valve of FIG. 29A in a closed configuration.

FIG. 29D is a cross sectional view of the duckbill check valve of FIG. 29A in an open configuration.

FIG. 30A is a perspective view of the connector of FIG. 27A, and a syringe, and a vial in an unassembled configuration.

FIG. 30B is a perspective view of the connector of FIG. 27A, and a syringe, and a vial in an assembled configuration.

FIG. 30C is a front view of the connector of FIG. 27A.

FIG. 31A is a cross sectional view of the connector of FIG. 27A, a vial, and a syringe as fluid is drawn from the vial, through the connector, and into the syringe.

FIG. 31B is a cross sectional view of the connector of FIG. 27A, a vial, and a syringe as fluid is driven from the syringe, through the connector, and into an IV bag.

FIG. 32A is a perspective view of another embodiment of a connector for transferring fluid.

FIG. 32B is another perspective view of the connector of FIG. 32A.

FIG. 33A is an exploded perspective view of the connector of FIG. 32A.

FIG. 33B is another exploded perspective view of the connector of FIG. 32A.

FIG. 34A is a cross sectional view of the connector of FIG. 32A, a vial, and a syringe as fluid is drawn from the vial, through the connector, and into the syringe.

FIG. 34B is a cross sectional view of the connector of FIG. 32A, a vial, and a syringe as fluid is driven from the syringe, through the connector, and into an IV bag.

FIG. 35A is a perspective view of another embodiment of a connector for transferring fluid.

FIG. 35B is another perspective view of the connector of FIG. 35A.

FIG. 36A is an exploded perspective view of the connector of FIG. 35A.

FIG. 36B is another exploded perspective view of the connector of FIG. 35A.

FIG. 37 is a perspective view of a check valve assembly that can be used with the connector of FIG. 35A.

FIG. 38A is a cross sectional view of the connector of FIG. 35A, a vial, and a syringe as fluid is drawn from the vial, through the connector, and into the syringe.

FIG. 38B is a cross sectional view of the connector of FIG. 35A, a vial, and a syringe as fluid is driven from the syringe, through the connector, and into an IV bag.

FIG. 39 is a perspective view of a system for transferring precise amounts of fluid.

FIG. 40 is a perspective view of a fluidics assembly for use with the system of FIG. 39.

FIG. 41 is an exploded perspective view of the fluidics assembly of FIG. 40.

FIG. 42 is an exploded perspective view of a vial adapter.

FIG. 43 is a cross sectional view of the vial adapter of FIG. 42.

5

FIG. 44 is a perspective view of a connector of the fluidics assembly of FIG. 40.

FIG. 45 is another perspective view of the connector of FIG. 44.

FIGS. 46-51 show various views of the connector of FIG. 44.

FIGS. 52-53 are exploded perspective views of the connector of FIG. 44.

FIGS. 54-55 are cross sectional views of the connector and syringe of the fluidics assembly of FIG. 40.

FIG. 56 is a perspective view of the IV bag assembly of the fluidics, system of FIG. 40.

FIG. 57 is an exploded perspective view of another sample embodiment of an IV bag assembly.

FIG. 58 is a perspective view of a top connector of the system of FIG. 39.

FIG. 59 is a perspective exploded view of the top connector of FIG. 58.

FIGS. 60-65 show various views of the top connector of FIG. 58.

FIGS. 66-71 show various views of the cassette of the top connector of FIG. 58.

FIGS. 72-77 show various views of the base member of the top connector of FIG. 58.

FIG. 78 is a cross sectional view of the second male connector of the connector of FIG. 44.

FIG. 79-81 are perspective views of the top connector that are cut and separated to illustrate the interior of the top connector.

FIG. 82 is a top-down view of the top connector and syringe of FIG. 81.

FIG. 83 is a side view of a tray attached to the top connector.

FIG. 84 is a side view of the tray and top connector in a disengaged configuration.

FIG. 85 is a flowchart showing an embodiment for priming the fluidics assembly of FIG. 40.

FIG. 86 is a flowchart showing an embodiment for transfer fluid.

FIG. 87 is a flowchart showing an example embodiment for replacing a vial during the transfer of fluid.

FIG. 88 is a perspective view of another example embodiment of a system for transferring fluid.

FIG. 89 is a perspective view of a top connector from a fluid transfer station of the system of FIG. 88.

FIG. 90 is a perspective view of the tray associated with the top connector of FIG. 89.

FIG. 91 is a perspective view of the top connector of FIG. 89 with the tray attached thereto in a first configuration.

FIG. 92 is a perspective view of the top connector of FIG. 89 with the tray attached thereto in a second configuration.

FIG. 93 is a split perspective view of the top connector of FIG. 89 and the tray.

FIG. 94 is a cross sectional view of the top connector of FIG. 89 and the tray.

FIG. 95 is a perspective view of the cassette from the top connector of FIG. 89.

FIG. 96 is a front view of the cassette of FIG. 95.

FIG. 97 is a cross sectional view of the connector shown in FIG. 88 with an outline of the cassette from FIG. 95.

FIG. 98 is a perspective view of another example embodiment of a connector for transferring fluid.

FIGS. 99-104 are cross sectional views of the target connector piece taken along the line 99-99 of FIG. 97 with the housing positioned as various different rotational positions.

FIG. 105 is a side view of another example embodiment of a connector that can be used to transfer fluid.

6

FIG. 106 is a cross sectional view of the target connector portion of the connector of FIG. 105.

FIG. 107 is a perspective view of another example embodiment of a connector that can be used to transfer fluid.

FIG. 108 is a cross sectional view of the target connector portion of the connector of FIG. 107 with the valve member in the closed position and an unobstructed light path.

FIG. 109 is a cross sectional view of the target connector portion of the connector of FIG. 107 with the valve member in the open position and an obstructed light path.

FIG. 110 is a cross sectional view of the target connector portion of the connector of FIG. 107 with the valve member in the closed position and an obstructed light path.

FIG. 111 is a cross sectional view of the target connector portion of the connector of FIG. 107 with the valve member in the open position and an unobstructed light path.

DETAILED DESCRIPTION OF SOME EXAMPLE EMBODIMENTS

The following detailed description is now directed to certain specific example embodiments of the disclosure. In this description, reference is made to the drawings wherein like parts are designated with like numerals throughout the description and the drawings.

In many circumstances fluid is transferred from a source container to a target container. In some instances, it can be desirable to transfer precise amounts of a fluid such as a medication into the target container. For example, in some embodiments a medication can be stored in a vial or other container, and a precise dosage amount of the medication can be extracted and transferred to a target device so that the dosage amount can be delivered to a patient. In some embodiments, fluid from multiple source containers can be combined, or compounded, into a single target container. For example, in some embodiments a mixture of medications can be created in the target container, or a concentrated medication can be combined with a diluent in the target container. To achieve the desired proportions of fluids, it can be desirable to precisely measure the amounts of fluids transferred into the target container. Also, precisely measuring the amount of fluid transferred from the source container to the target container can reduce the amount of fluid wasted (e.g., when more fluid than necessary is withdrawn from the source container). Reduction of waste is desirable because in some instances the fluid being transferred can be expensive.

Some embodiments disclosed herein provide a fluid transfer device for transferring precise amounts of fluid from one or more source containers into one or more target containers.

In some embodiments, it can be desirable to transfer fluids from a source container to a target container using a sealed system. In some embodiments, exposing the fluid to ambient air can allow contaminants to enter the fluid or cause an undesirable reaction with the fluid. Some medications (e.g., chemotherapy medications) can be harmful to a healthy individual. Therefore, it can be desirable to prevent or reduce exposure of the fluid being transferred to the ambient air or area outside the fluid transfer system. In some embodiments, a fluid transfer system that prevents or reduces exposure of the fluid to the area outside the fluid transfer system can render other expensive equipment (e.g., a clean room) unnecessary, thereby reducing the cost associated with transferring the fluids.

Some embodiments disclosed herein provide a fluid transfer device for transferring fluid while preventing, reducing, or minimizing the amount of contact the fluid has with the ambient air or area outside the fluid transfer system.

FIG. 1 schematically shows an embodiment of an automated fluid transfer system 100. The system 100 can include a housing 102 enclosing a controller 104 and a memory module 106. The system 100 can also include a user interface 108, which can be, for example, external to the housing 102. The user interface 108 can also be integrated into the housing 102 in some cases. The user interface 108 can include, for example, a display, a keypad, and/or a touch screen display. The user interface 108 can be configured to receive instructions from the user, for example, regarding the amounts of fluid to be transferred and the types of fluids to be transferred. The user interface can also be configured to provide information to the user, such as error messages, alerts, or instructions (e.g., to replace an empty vial). The system 100 can also include a bar code scanner 110 in communication with the controller 104. Although in the embodiment shown, the controller 104 and memory module 106 are contained within the housing 102, a variety of other configurations are possible. For example, controller 104 can be external to the housing 102, and can be, for example contained within a second housing which also contains the user interface 108. In some embodiments, the system 100 can include a communication interface 105 configured to receive information (e.g., instructions) from a remote source such as a terminal or an automated management system, etc. In some embodiments, the communication interface can also send information (e.g., results or alerts) to the remote source. In some embodiments, the system 100 does not include a communication interface 105 and does not communicate with a remote source.

The system 100 can include multiple transfer stations 112a-c. In the embodiment shown, the system 100 includes three transfer stations 112a-c, but a different number of transfer stations can be used. For example, in some embodiments, the system may include a single transfer station. In other embodiments, the system may include two, four, five, six, seven, eight, or more transfer stations depending on the number of different fluid types the system is designed to handle and the amount of fluid to be transferred.

Each transfer station 112a-c can include a fluid source container 114a-c, which can be, for example, a medical vial or other suitable container such as a bag, a bottle, or a vat, etc. Although many embodiments disclosed herein discuss using a vial as the source container, it will be understood the other containers can be used even when not specifically mentioned. In some embodiments, each of the source containers 114a-c can contain a unique fluid, providing a variety of fluids that the user can select for transfer. In other embodiments, two or more of the source containers 114a-c can contain the same fluid. In some embodiments, the source containers 114a-c include bar codes that identify the types of fluid contained therein. The bar codes can be scanned by the scanner 110 so that the identities of the fluids contained by source containers 114a-c can be stored within memory module 106. In some embodiments, the fluid transfer stations 112a-c are configured to transfer precise amounts of fluid from source containers 114a-c to target containers 116a-c, which can be, for example IV bags. It will be understood that in various embodiments described herein, a different type of target connector or destination container can be used instead of an IV bag (e.g., a syringe, a bottle, a vial, etc.) even when not specifically mentioned. In some embodiments the fluid can first be transferred from source containers 114a-c to intermediate measuring containers 118a-c so that a precise amount of fluid can be measured. The intermediate measuring containers 118a-c can be, for example, syringes. After being measured, the fluid can be transferred from intermediate measuring containers 118a-c to the target containers 116a-c. In some

embodiments, one or more of the transfer stations 112a-c can include one or more pairs of male and female fluid connectors configured to be attached to each other to selectively permit the passage of fluid. When fluid transfer is completed, the connectors can be detached or disconnected. In some embodiments, the connectors can be configured to automatically close. The fluid module can be removed while retaining substantially entirely or entirely all of the remaining interior fluid within the respective connectors and the rest of the fluid module, thus permitting the transfer to occur in a substantially entirely or entirely closed system, thereby diminishing the risk of damage caused by liquid or vapor leakage from the fluid module after disconnection and from the fluid source and the fluid destination after disconnection.

In some embodiments, the system 100 can be configured to be compatible with a variety of sizes of syringes. For example, larger volume syringes can be used to transfer larger volumes of fluid in shorter amounts of time. Smaller volume syringes can be used to increase the accuracy and precision with which amounts of fluid can be transferred. In some embodiments, the syringes can include a bar code which identifies the volume of the syringe. The bar code can be scanned by a bar code scanner 110, so that the sizes of the syringes used by the different transfer stations 112a-c can be stored within memory module 106 for use by the controller 104.

In some embodiments, connectors 120a-c connect the source containers 114a-c, the intermediate containers 118a-c, and the target containers 116a-c. In some embodiments, the connectors 120a-c can include first check valves (not shown) configured to allow fluid to flow from the source containers 114a-c into the connector 120a-c, and block fluid from flowing connector 120a-c into the source containers 114a-c, as shown by single-headed arrows. The connectors 120a-c can also include second check valves (not shown) configured to allow fluid to flow from connectors 120a-c into target containers 116a-c, but block fluid from flowing from target containers 116a-c into connectors 120a-c, as shown by single-headed arrows. In some embodiments, the connectors 120a-c can be in two-way fluid communication with the intermediate containers 118a-c, as shown by double-headed arrows.

In some embodiments, the system 100 can include mounting modules 122a-c for mounting the transfer stations 112a-c onto the housing 102. For example, in some embodiments the mounting modules 122a-c can be configured to securely receive intermediate measuring containers 118a-c as shown in FIG. 1. The system 100 can also include motors 124a-c, which can be for example, contained within housing 102. The motors 124a-c can be configured to actuate the plungers on the syringes 118a-c to draw fluid into the syringes and to dispel fluid therefrom. The motors 124a-c can be in communication with the controller 104, and can receive actuation instructions from the controller 104.

In some embodiments, the system can include fluid detectors 126a-c configured to detect a presence or absence of fluid in connectors 120a-c. The fluid detectors 126a-c can be in communication with the controller 104 so that when the detectors 126a-c detect an absence of fluid in connectors 120a-c, indicating that source fluid containers 114a-c have run dry, they can send a signal to controller 104 that a source container 114a-c needs to be replaced. The fluid detectors 126a-c can be for example an infrared LED and photo detector, or other type of electronic eye, as will be discussed in more detail below. In the embodiment shown, fluid detectors 126a-c are shown connected to connectors 128a-c, but other

configurations are possible. For example, fluid detectors **126a-c** can be connected to fluid source containers **114a-c** themselves.

In some embodiments, the system **100** can include compatibility mechanisms **127a-c** for ensuring that an approved connector **120a-c** has been placed in communication with the system **100** to ensure the accuracy of the amount of fluid transferred. The compatibility mechanisms **127a-c** can be, for example, a specifically shaped mounting feature configured to correspond to a portion of the connector **120a-c**.

In some embodiments, the system **100** can include source adapters **129a-c** configured to receive the source containers **114a-c** and removably connect to the connectors **120a-c**. Thus, when a source container **114a-c** runs out of fluid, the empty source container **114a-c** and its corresponding adapter **129a-c** can be removed and replaced without removing the associated connector **120a-c** from the system **100**. In some embodiments, source adapters **129a-c** can be omitted, and the source containers **114a-c** can be directly received by the connectors **120a-c**.

In some embodiments the system **100** can include sensors **128a-c** for detecting the presence of target containers **116a-c**. Sensors **128a-c** can be in communication with the controller **104** so as to prevent the system **100** from attempting to transfer fluid when no target container **116a-c** is connected. A variety of sensor types can be used for sensors **128a-c**. For example, sensors **128a-c** can be weight sensors or infrared sensors or other form of electronic eye. In some embodiments, weight sensors **128a-c** can also be used to measure the weight of the target containers **116a-c** after fluid has been transferred. The final weight of a target container **116a-c** can be compared to an expected weight by the controller **104** to confirm that the proper amount of fluid was transferred into the target container **116a-c**. Sensors **128a-c** can be a variety of other sensor types, for example sensor pads or other sensor types able to detect the presence of target containers **116a-c**.

FIG. 2 schematically illustrates a system **200** for automated precise transfer of fluids. System **200** can be the same as or similar to the system **100** in some regards. Some features shown in FIG. 1, such as the adapters **129a-c** and compatibility mechanisms **127a-c**, are not shown specifically in the system **200**, but it will be understood that system **200** can include corresponding features. The system **200** can include a housing **202**, a controller **204**, a memory **206**, a user interface **208**, a scanner **210**, and a communication interface **205**, similar to those describe above in connection with the system **100**. System **100** is configured to transfer individual fluids from the source containers **114a-c** to target containers **116a-c**. System **200**, on the other hand, is configured to transfer and combine fluids from source containers **214a-c** into a common target container **216**. Thus, system **200** can be used for compounding mixtures of fluids. In some embodiments, a single system can be configured both for compounding mixtures of fluids and for the transfer of individual fluids from a single-source container to a single-target container. For example, a system containing six fluid transfer stations can be configured so that transfer stations **1-3** are dedicated to compounding mixtures of fluids into a single common target container, while fluid transfer stations **4-6** can be configured to each transfer fluid from a single source container to a single target container. Other configurations are possible. In the embodiment shown in FIG. 2, the system **200** can include sensors **228a-c** for detecting whether or not the connectors **220a-c** are connected to the common target container **216**. The system **200** can also include a sensor **229** for detecting the presence of the common target container **216**. In some embodiments, the sensor **229** can measure the weight of the common target container

216 and can report the weight to the controller **104**. The controller **104** is then able to compare the final weight of the common target container **216** with an expected weight to confirm that the common target container **152** was filled with the correct amount of fluids.

FIGS. 3A and 3B show a subsystem, or fluidics assembly, **300** for transferring precise amounts of fluid from a medical vial **314** to an IV bag **316**. FIG. 3A is a perspective view of subsystem **300**, and FIG. 3B is an exploded perspective view of subsystem **300**. The subsystem **300** can include a syringe **318** for measuring precise amounts of fluid to be transferred. In some embodiments, the system includes an IV bag assembly **330**. The IV bag assembly **330** can include the IV bag **316**, a connector **332**, and a piece of tubing **334** connecting the IV bag **316** to the connector **332**. The connector **332** can be, for example, a female medical connector. The connector **332** illustrated in FIGS. 3A-B is a version of the Clave® connector manufactured by ICU Medical, Inc., of San Clemente, Calif. Various embodiments of a connector of this type are described in U.S. Pat. No. 5,685,866 (the “866 Patent”), the entirety of which is incorporated herein by reference. The subsystem **300** can also include a connector **320**, for interconnecting the vial **314**, the syringe **318**, and the IV bag assembly **330**.

Turning now to FIGS. 4A and 4B, FIG. 4A shows an exploded perspective view of a fluid transfer module in the form of connector **320**, and FIG. 4B shows a cross-sectional view of the connector **320**. The connector **320** can include a first interface or source connector portion **336** configured to provide fluid communication between the connector **320** and the vial **314**, a second interface of target connector portion **338** configured to provide fluid communication between the connector **320** and the IV bag assembly **330**, and an intermediate connector portion **340** configured to provide fluid communication between the connector **320** and the syringe **318**. The connector can also include a main body **342**. In the embodiment shown in FIGS. 4A-B, the intermediate connector portion **340** is integrally formed as part of the main body **342**.

In some embodiments, the connector **320** can be a T-connector. In the embodiment shown, the fluid path leading to the IV bag assembly **330** is substantially perpendicular to the fluid path between the vial **314** and the syringe **318**. A variety of other configurations are possible. For example, the fluid pathways can be arranged to intersect at an oblique angle.

In some embodiments, the source connector portion **336** includes a female connector portion **344** having a slightly tapered internal surface. The main body **342** of the connector can have a corresponding male connector portion **346** having a similarly tapered outer surface. The female connector portion **344** and male connector portion **346** can be configured such that when the male connector portion **346** is fully inserted into the female connector portion **344** (i.e., the tapered surfaces prevents further insertion), a chamber **348** is defined between the end of the male connector portion **346** and the base of the female connector portion **344**. The male connector portion **346** can be secured to the female connector portion **344** by applying a plastic welding adhesive (such as Dichloromethane) to the outer surface of the male connector portion **346** and/or to the inner surface of the female connector portion **344** before insertion. The Dichloromethane can chemically weld the outer surface of the male connector portion **346** to the inner surface of the female connector portion **344**. Other methods can be used to connect the male connector portion **346** to the female connector portion **344**, such as sonic welding, threading, adhesives, etc. In some embodiments, the connection between the main body **342** and

11

the source connector portion 336 is hermetically sealed, and in some embodiments includes a sealing member (not shown), such as an O-ring, to provide the hermetic seal.

In some embodiments, the target connector portion 338 can be similarly attached to the main body 342. The main body 342 can include a female connector portion 350 with a tapered inner surface, and the target connector portion 338 can include a male connector portion 352 with a tapered outer surface. When the male connector portion 352 is inserted fully into the female connector portion 350 (i.e., the tapered surfaces prevent further insertion), a chamber 354 is defined between the end of the male connector portion 352 and the base of the female connector portion 350. The connector portions 350, 352 can be secured to one another using Dichloromethane or any of the other methods discussed above. In some embodiments, the connection between the main body 342 and the target connector portion 338 is hermetically sealed, and in some embodiments, the connection can include a sealing member.

The connector 320 can include a source check valve 356 disposed inside the chamber 348. The check valve 356 can be configured to allow fluid to flow from the vial 314 into the connector 320, but block fluid from flowing from the connector 320 into the vial 314. The connector can also include a target check valve 358 disposed inside chamber 354. Check valve 358 can be configured to allow fluid to flow from the connector 320 into the IV bag assembly, but blocks fluid from flowing from the IV bag assembly into the connector 320. The check valves 356, 358 will be discussed in greater detail below.

The main body 342 can be constructed from a variety of materials. The main body 342 can be constructed from a rigid material such as polycarbonate or other polymeric materials. In some embodiments, at least a portion of the main body 342 can be formed from a substantially transparent material as discussed below.

FIG. 5A shows a perspective view of the source connector portion 336 and vial 314 in an unengaged configuration. FIG. 5B is another perspective view of the source connector portion 336 and vial 314, also in an unengaged configuration. FIG. 5C is a cross-sectional view of the source connector portion 336 and vial 314 in an engaged configuration. FIG. 5D is a cross-sectional view of the source connector portion 336 and vial 314 after a portion of the fluid has been withdrawn from the vial 314. Although FIGS. 5A-5D shown the source connector portion 336 of the connector 320 separated from the remainder of the connector 320 for simplicity, it should be understood that the source connector portion 336 can be connected to the remainder of the connector 320 when in use.

With reference now to FIGS. 5A-D, the vial 314 can comprise any suitable container for storing medical fluids, and can be for example a medical vial such as those produced by Abbott Laboratories of Abbott Park, Ill. In some embodiments, the vial 314 includes a body 357 and a cap 359. In some instances, the vial 314 can be configured to be hermetically sealed. The body 357 can comprise a rigid substantially impervious material such as plastic or glass. In some embodiments the cap 359 includes a septum 360 and casing 362. The septum 360 can be made of an elastomeric material capable of deforming in such a way that when punctured by an item, it forms a substantially airtight seal around that item. For example, in some instances the septum 360 comprises silicone rubber or butyl rubber. The casing 362 can surround the septum 360 and can be made from any suitable material for sealing the vial 314. In some instances, the casing 362 comprises a metal that is crimped around the septum 360 and an

12

end portion of the vial body 357 in order to form an airtight seal between the septum 360 and the vial body 357. In some embodiments, casing 362 can include a substantially flat mounting surface 364. The vial 314 can include a fluid 366, such as a medical fluid (e.g., a chemotherapy drug) contained within its internal volume. The vial 314 can also include a relatively small amount of sterilized air 368 also contained within the internal volume.

The source connector portion 336 can include a piercing member 370 which can comprise a sheath 372 and a pointed tip 374. The sheath 372 can be cylindrical in shape, or it can be a variety of other suitable shapes. For example, in some embodiments, the sheath 372 can be generally conical in shape and taper toward the pointed tip 374. The piercing member 370 can comprise a rigid material such as metal or plastic, suitable for insertion through the septum 360, such as a polycarbonate plastic. In some instances the pointed tip 374 is separable from the sheath 372. In other embodiments, the pointed tip 374 and sheath 372 can be integrally formed or permanently joined. The pointed tip 374 can be configured to facilitate piercing of the septum 360. The source connector portion 336 can also include a cap connector 376 configured to secure the source connector portion 336 to the vial 314. In some embodiments, the cap connector 376 can include an adhesive 378, such as a double-sided tape, disposed on the surface of the cap connector 376. A removable covering 380 (shown partially peeled away in FIG. 5B) can be disposed over the adhesive 378 until it is ready to be used. The vial 314 can be secured to the cap connector 376 by removing the covering 380 from the adhesive 378 and pressing the vial 314 down onto the source connector portion 336 so that the piercing member 370 pierces the septum 360 and the mounting surface 364 comes into contact with the adhesive 378. A variety of other connection types can be used to secure the vial 314 to the source connection portion 336 of the connector 220.

In some embodiments, the source connector portion 336 can be configured to automatically equalize pressure within the vial 314 as fluid 366 is withdrawn. For example, the source connector portion 336 can be a version of the Genie® closed vial access device manufactured by ICU Medical, Inc. of San Clemente, Calif. Certain embodiments of closed vial access devices of this type are disclosed in U.S. Provisional Patent Application No. 61/090,561 (the “’561 Application”), the entirety of which is herein incorporated by reference. For example, the ’561 Application discloses other methods by which the vial 314 can be connected to the source connector portion 336.

In some embodiments, the source connection portion 336 can include a fluid extraction channel 382. The fluid extraction channel 382 can include an upper portion 384 that extends from an extraction aperture 383 formed in the side wall of the piercing member 370 through a portion of the piercing member 370. The fluid extraction channel 382 can also include a lower portion 386 that extends through the female connection portion 344. In certain embodiments, the lower portion 386 can be wider than the upper portion 384, defining a shoulder 388 at the transition from the lower portion 386 to the upper portion 384.

In some embodiments, the sheath 372 can be hollow defining a regulator channel 390 that extends through the sheath 372 and through the cap connector 376 to a regulator aperture 392 formed on a position of the source connector portion 344 that remains exposed to the ambient air when the vial 314 is secured to the source connector portion 336. In some embodiments, a bag 394 can be enclosed within the regulator channel 390. The bag can define an inner volume 395 that is in fluid

13

communication with the regulator channel 390. In some embodiments, the bag can include a connection region 396 that forms an airtight seal with the walls of the regulator channel 390 so that air cannot move past the connection region 396 unless it enters the inner volume 395 of the bag 394. In some embodiments, the connection region 396 of the bag 394 can be secured to the sheath 372 by an adhesive, or by any other suitable manner.

The bag 394 can be folded up inside the regulator channel 390 so that it occupies a relatively small volume compared to its unfolded state. The bag 394 can be configured to be able to fill all, or a substantial portion, of the internal volume of the vial 314. In some embodiments, the bag 394 can comprise an elastomeric material, such as Mylar®, polyester, polyethylene, polypropylene, saran, latex rubber, polyisoprene, silicone rubber, polyurethane, and latex-free silicone that can allow the bag 394 to unfold, expand, and/or contract. In some embodiments, the bag 394 can comprise a non-expandable material that is flexible enough to allow the bag to unfold. In some circumstances, the bag 394 can comprise a material that is impervious to liquid and air and inert with respect to the fluid 366.

FIG. 5C illustrates an embodiment of the source connector portion 336 coupled to the vial 314 at a stage before any of the fluid 366 is extracted. By comparison, FIG. 5D illustrates an embodiment of the source connector portion 336 coupled to the vial 314 at a stage with the bag 394 deployed after some of the fluid 366 has been extracted. Although not shown in FIGS. 5C and 5D, the fluid extraction channel 382 of the source connector portion 336 can be in fluid communication with the syringe 318 or other medical instrument capable of creating a negative pressure to extract fluid 366 from the vial 314. In some circumstances, a volume of the fluid 366 can be withdrawn from the vial 314 by the syringe causing the pressure within the vial 314 to drop. The reduced pressure in the vial can cause the tip 374 to disengage from the sheath 372, so that the bag 394 is free to emerge from the sheath 372. As the fluid 366 flows out of the vial 314 and toward the syringe 318, ambient air flows into the inner volume 395 of the bag 394 by way of the regulator channel 390 and the regulator aperture 392. In some circumstances the inner volume 395 of the bag 394 expands (by the bag unfolding and/or expanding) to compensate for the reduced pressure inside the vial 314.

Thus, the source connector portion 336 can be configured to allow the fluid 366 to be withdrawn from the vial 314 while regulating the pressure within the vial 314. In some embodiments, the source connector portion 336 maintains a substantially constant pressure within the vial 314 as the fluid 366 is withdrawn therefrom. In some embodiments, the pressure within the vial 314 changes by no more than about 1-5 psi as the fluid 366 is withdrawn. The '561 Application discloses additional details and various alternatives that can be applied to the source connector portion 336 and vial 314.

FIG. 6A shows a perspective view of the target connector portion 388. FIG. 6B is an exploded perspective view of the target connector portion 388. FIG. 6C shows a top view of a housing portion of the target connector portion 388. FIG. 6D shows a cross-sectional view of the target connector portion 388 and the female connector 332 in an unengaged configuration. FIG. 6E shows a cross-sectional view of the target connector portion 388 and the female connector 332 in an engaged configuration. Although the target connector portion 388 is shown separated from the remainder of the connector 320 in FIGS. 6A-6E, it should be understood that the target connector portion 388 can be connected to the remainder of the connector 320 when in use.

14

With reference now to FIGS. 6A-6E, the target connector portion 388 of the connector 320 can be a closeable male luer connector that is configured to prevent fluid from escaping from or entering into the connector when it is not engaged with a corresponding female connector, but allow fluid to flow when it is engaged with a corresponding female connector 332. In the embodiments shown, the target connector portion 388 can be a version of the Spiros® closeable male connector manufactured by ICU Medical, Inc., of San Clemente, Calif. Various embodiments of connectors of this type are described in U.S. Patent Publication No. 2008/0287920 (the "920 Publication"), the entirety of which is incorporated herein by reference. Although the embodiments illustrated in FIGS. 6A-6E show the connector 332 as being a female connector and the target connector portion 388 as being a male connector, it should be noted that other configurations are possible. For example, the connector 332 can be a male connector while the target connector portion 388 can be a female connector. In some embodiments, a substantially entirely or entirely closed system can be achieved, at least in part, by providing corresponding automatically closeable male and female connectors at various (or all) connection points within the fluid transfer system 100, thereby causing the stationary fluid to substantially entirely remain within the fluid source, the fluid module, and the fluid target, respectively, upon disconnection and to not generally leak or vaporize outside of the system. For example, in some embodiments, corresponding pairs of automatically closing connectors (e.g., male and female connectors) can be provided at the interfaces between the fluid source and the fluid module, the fluid module and the intermediate container, and/or the fluid module and the destination or target container.

The target connector portion 388 can include a housing 398, a valve member 400, a resilient member 402, a sealing ring 404, an end cap 406, and an O-ring 407. The housing 398 can be generally tubular in shape, and can include a passageway 408 that extends axially through the housing. As illustrated, the passageway 408 includes apertures on each side of the connector. The housing 398 can include a male luer tip 410 that connects to the rest of the housing 398 at a base 412. The luer tip 410 can be generally tubular in shape so that a portion of the passageway 408 is defined therein, and the luer tip 410 can include a hole 414 at its end providing access to the passageway 408. In some embodiments, the luer tip 410 includes a shelf 416 that extends radially inwardly toward the axis of the passageway 408. The shelf 416 can be located adjacent to the hole 414, so that the passageway 408 is narrowed at the end of the luer tip 410. In some embodiments, the surface of the shelf 416 that faces radially inwardly is tapered so that the passageway 408 is narrowest immediately adjacent to the hole 414. In some circumstances, the shelf 416 can be configured to seal the passageway when a portion of the valve member 400 is abutted against it. As illustrated, in some embodiments, connectors can be used to substantially entirely prevent fluid therein to leak, vaporize, or otherwise escape through apertures in the fluid pathway when the connectors are closed.

The luer tip 410 can be surrounded by a shroud 418. In some embodiments, the luer tip 410 extends some distance beyond the edge 420 of the shroud. The shroud 418 can include inner threads 422 on its interior surface. The inner threads 422 can be used for securing a female connector 332. The shroud can include an indented portion 424 that has a smaller outer diameter than the other portions of the housing. The indented portion 424 can be configured to engage a portion of the resilient member 402.

15

The housing 398 can include two wall sections 426a, 426b separated by two gaps 428a, 428b. The gaps 428a, 428b can be configured to receive portions of the resilient member 402. The wall sections 426a, 426b can be configured to engage the end cap 406.

In some embodiments, the housing 398 includes a middle portion 430 located substantially between the wall sections 426a, 426b, and connected to the wall sections 426a, 426b near the gaps 428a, 428b. In some embodiments, holes 432a, 432b are defined between the middle portion 430 and the wall sections 426a, 426b (as shown in FIG. 6C). In some embodiments, the luer tip 410 connects to the middle portion 430 at its base 412. In some embodiments, the middle portion defines a portion of the passageway 408 therein. In some embodiments, portions 434 of the outer surface of the middle portion 430 are exposed by the gaps 428a, 428b. The portions 434 can include notches 436a, 436b and through-holes 438a, 438b. The notches 436a, 436b can be generally rectangular in shape, and can be tapered such that the notches 436a, 436b are narrower near their bases than near their surfaces. The through-holes 438a, 438b can also be generally rectangular in shape.

The housing 398 can be constructed from a variety of materials. The housing 398 can be constructed from a rigid material such as polycarbonate or other polymeric materials. In some embodiments, the housing 398 can be constructed from a hydrophobic material such as Bayer Makrolon, or any other suitable material. In some embodiments, the housing 398 can be formed from a substantially transparent material.

The valve member 400 can include a fluid passageway 440 extending axially from an opening formed in a base portion 444 and into a tube 446. In some embodiments, the passageway 440 can be wider in the base portion 444 than in the tube 446. In some embodiments, the tube 446 includes a narrowed tip 448. In some embodiments, the tip 448 can have a tapered outer surface. The tip 448 can be tapered to substantially the same degree as the radially inwardly facing surface of the shelf 416 and can be sized so that the tip 448 can form a fluid seal with the shelf 416 when abutted against it. In some embodiments, the tip 448 can be made from a flexible or compressible material, such as silicone rubber to facilitate formation of the fluid seal between the tip 448 and the shelf 416. In some embodiments, the tube can include one or more holes 450 for providing access to the fluid passageway 440. The holes 450 can be formed, for example, in the tip 448 of the tube 446.

In some embodiments, the valve member 400 can include two struts 452a, 452b extending out from the base 444 and positioned on either side of tube 446, so that an open space is defined on either side of the tube. In some embodiments, the tube 446 can extend axially past the ends of the struts 452a, 452b.

The base 444 of the valve member 400 can include a plurality of protrusions 454 extending radially outwardly from its external surface. In some embodiments, the protrusions 454 can be positioned so as to define two channels 456a, 456b therebetween. In some embodiments, the protrusions 454 do not extend across the full length of the base 444, leaving a lower portion 458 of the base 444 that has a substantially smooth outer surface.

The valve member 400 can be constructed from a variety of materials, such as polycarbonate or other polymeric materials. In some embodiments, the valve member 400 can be constructed from the same material as the housing 398. In some embodiments, the valve member 400 and housing 398 can be constructed from different materials. In some embodiments, the valve member 400 can be constructed from mul-

16

tle materials or from multiple pieces. For example, the tip 448 can be constructed from a material that is more flexible than the remainder of the valve member 400. In some embodiments, the valve member 400 can be formed from a substantially opaque material.

The resilient member 402 can include a first ring 460 and a second ring 462 connected to each other by elastic members 464a, 464b. The elastic members 464a, 464b can be made from an elastic material that exerts a restoring force when stretched, such as silicon rubber. Thus, if the rings 460, 462 are pulled apart, the elastic members 464a, 464b function to restore the rings 460, 462 to their relaxed configuration. In some embodiments, the rings 460, 462 are also constructed from an elastic material, such as the same material used to form the elastic members 464a, 464b. In some embodiments, the second ring 462 can have a greater diameter than the first ring 460. In some embodiments, the second ring 462 can have a tapered outer surface so that the end of the second ring 462 that is closest to the first ring 460 is wider than the end of the second ring 462 that is furthest from the first ring 460.

The sealing ring 404 can be generally cylindrical in shape, and can have a bore 466 extending axially therethrough. The sealing ring 404 can have a cylindrical body section 468 and an O-ring 470 located at one end of the body section 468. In some embodiments, the thickest portion of the O-ring 470 can be thicker than the body section 468 so that the thickest portion of the O-ring 470 extends radially inwardly toward the axis of the bore 466 a distance past the inner surface of the body section 468. Thus, the bore 466 can be narrower at the thickest part of the O-ring 470 than in the body section 468. In some embodiments, the thickest portion of the O-ring 470 also extends radially outwardly a distance past the outer surface of the body section 468. The sealing ring 404 can include two protrusions 472a, 472b that extend radially outwardly from the body section 468. In some embodiments, the protrusions 472a, 472b can be generally rectangular in shape.

The sealing ring 404 can be constructed from a variety of materials. In some embodiments, the sealing ring 404 can be constructed from a deformable or elastic material such as a silicone rubber. In some embodiments, the sealing ring 404 can be constructed from the same material used for form the resilient member 402. In some embodiments, the sealing ring 404 can be constructed from a material capable of forming a fluid seal against a rigid plastic or other rigid polymeric material.

The end cap 406 can include a first end cap member 405 and a second end cap member 409. The second end cap member 409 can include a male connector 352, a plunger 474, and a disk portion 476 located between the male connector 352 and the plunger 474. The second end cap member 409 can have a fluid passageway 478 axially positioned therein. In some embodiments, the plunger 474 can be generally tubular in shape. In some embodiments, the outer surface of the plunger 474 includes an indented region 480, which can be configured to receive the O-ring 407 therein. The O-ring 407 can be constructed from an elastic material such as silicone rubber so that it can be stretched over the edge 482 of the plunger 474 and be seated in the indented region 480. In some embodiments, the O-ring 407 can be constructed from the same material as the resilient member 402 and/or the sealing ring 404. In some embodiments, the O-ring 407 can be sized so that when seated in the indented region 480, the thickest portion of the O-ring 407 extends radially outwardly a distance past the outer surface of the plunger 474.

In some embodiments, the passageway 478 can have a substantially constant width throughout the second end cap member 409. In some embodiments, the passageway 478 can

be tapered so that it is wider in the male connector 352 than in the plunger 474. In some embodiments, the passageway 478 can narrow near the end of the plunger 474, for example, to accommodate the indented region 480.

The first end cap member 405 can be generally frustoconical in shape and can have a central opening 471 therein. When assembled, the plunger 474 can extend through the central opening 471. A ridge 473 can extend inward into the central opening 471. The ridge 473 can be received into a channel 475 formed between the base of the plunger 474 and the disk portion 476 on the second end cap member 409 to secure the first end cap member 405 to the second end cap member 409. The ridge 473 and corresponding channel 475 can allow the first end cap member 405 to rotate about a longitudinal axis with respect to the second end cap member 409. Thus, the first end cap member 405 and the second end cap member 409 can join to form the end cap 406.

The valve end cap 406 can be constructed from a variety of materials, such as polycarbonate or other rigid polymeric materials. In some embodiments, the end cap 406 can be constructed from the same material as the housing 398 and/or the valve member 400.

In some embodiments, the end cap 406 can be constructed from a different material than the valve member 400 and/or the housing 398. The first end cap member 405 can be formed from the same material as the second end cap member 409, or different materials can be used. In some embodiments, the first end cap member 405 or the second end cap member 409 or both can be substantially transparent.

Certain interconnections between various parts of the target connector portion 338 will now be discussed in further detail. The sealing ring 404 can be positioned inside the middle portion 430 of the housing 398. The protrusions 472a, 472b can be sized and positioned so that they engage the through-holes 438a, 438b. Thus, the sealing ring 404 can be secured to the housing 398 so that it does not rotate or move axially with respect to the tube 446.

The valve member 400 can be slidably inserted into the housing 398 so that the tube 446 enters the passageway 408. The narrowed tip 448 of the tube 446 can pass through the bore 466 of the sealing ring 404 and into the male luer tip 410 until it abuts against the shelf 416. The tube 446 can have a width that substantially fills the bore 446 and presses against the O-ring 470 portion of the sealing ring 404 to form a fluid seal therebetween. The struts 452a, 452b can pass through the holes 432a, 432b in the housing 398 respectively, so that the struts 452a, 452b are positioned between the male luer tip 410 and the shroud 418.

The resilient member 402 can function to bias the valve member 400 against the housing 398. The first ring 460 can fit onto the lower portion 458 of the base 444 of the valve member 400, so that a surface of the ring 460 abuts against the protrusions 454. The second ring 462 can fit into the indented portion 424 of the housing. The elastic members 464a, 464b can be positioned in the channels 456a, 456b respectively, and can pass through the respective gaps 428a, 428b between the wall sections 426a, 426b of the housing 398.

The O-ring 407 can be seated onto the indented region 480 of the end cap 406, as discussed above, and the plunger 474 can be slidably inserted at least partially into the passageway 440 of the valve member. In some embodiments, the thickest portion of the O-ring 407 can be wider than the portion of the passageway 440 formed in the base 444 of the valve member 400, so that the O-ring 407 forms a fluid seal against the inner surface of the passageway 440. The plunger 474 can be inserted into the valve member 400 until the disk portion 476

of the end cap 406 comes into contact with the ends of the wall sections 426a, 426b of the housing 398.

In some embodiments, the wall sections 426a, 426b can be secured to the top surface 477 of the first end cap member 405 by sonic welding, snap fit structures (not shown), a pressure or friction fitting, or other suitable connection type. As mentioned above, the first end cap member 405 can be secured to the second end cap member 409 in a manner that allows the first end cap member 405 to rotate relative to the second end cap member 409. Thus, once the target connector portion 338 is assembled, the housing 398, sealing ring 404, resilient member 402, valve member 400, and first end cap member 405 can rotate relative to the second end cap member 409 about the longitudinal axis.

With reference now to FIGS. 6D-6E, the target connector portion 338 can be configured to engage a female connector 332. A variety of types of female connectors 332 can be used. The female connector 332 shown is a closable female luer connector that includes a housing 490, a spike 492, a base 494, and a resilient seal element 496. A fluid passageway 498 can pass through the base 494 and through the spike 492. The spike 492 can include one or more holes 500 providing fluid communication between the passageway 498 and the area outside the spike 492. The seal element 496 can be shaped and positioned to substantially surround the spike 492. The seal element 496 can include a closable aperture 502 or slit that can open to allow the tip of the spike 492 to pass through then end of the seal element 496 when the seal element 496 is compressed (as shown in FIG. 6E). The housing can include external threads 504 configured to engage the inner threads 422 on the housing 398 of the target connector portion 338. An end of the tubing 334 can be connected to the end of the female connector 332 by an adhesive, clamp, friction or pressure fitting, or other suitable manner to form a fluid tight connection.

As discussed above, in some embodiments, the housing 398, sealing ring 404, resilient member 402, valve member 400, and first end cap member 405 can rotate about the longitudinal axis with respect to the second end cap member 409. Thus, as the female connector 332 of the IV bag assembly is attached to the target connector portion 338, the female connector 332 can be held still while the housing 398 of the target connector portion 338 can rotate causing the threads 504, 422 to engage. Because the female connector 332 is not required to rotate during engagement and disengagement with the target connector portion 338, the tubing 334 can avoid being twisted or kinked and the user is not required to twist the IV Bag to accommodate rotation of the female connector 332. Embodiments of the connectors with this rotational capability are disclosed in greater detail in the '920 Publication incorporated by reference herein in its entirety.

When not engaged with the female connector 332 (as shown in FIG. 6D), the target connector portion 338 can be sealed. In some embodiments, fluid can enter the target connector portion 338 at the male connector 352 and pass through the passageway 478 of the end cap 406, through the passageway 440 of the valve member 400, through the holes 450, and into the portion of the passageway 408 defined by the male luer tip 410. But the fluid seal created by the tip 448 of the valve member 400 pressing against the shelf 416 of the male luer tip 410 prevents the fluid from exiting the target connector portion 338. In some embodiments, an increase in pressure, such as when additional fluid is forced into the target connector portion 338, causes the tip 448 to press more firmly against the shelf 416, thereby improving the fluid seal.

When the target connector portion 338 is engaged with the female connector 332 (as shown in FIG. 6E), the external

threads 504 of the female luer connector 332 can engage the inner threads 422 on the shroud 418, securing the female connector 332 to the target connector portion 338. The edge of the male luer tip 410 can press against and compress the resilient seal element 496 so that the spike 492 passes through the aperture 502 until the holes 500 are exposed. The end of the housing 490 of the female luer connector 332 can enter the space between the male luer tip 410 and the shroud 418 until it contacts the struts 452a, 452b. As the female luer connector 332 further engages the target connector portion 338, it can push on the struts 452a, 452b causing the entire valve member 400 to retract. As the valve member 400 retracts, the elastic members 464a, 464b of the resilient member 402 stretch. When the valve member 400 retracts, the tip 448 disengages from the shelf 416, breaking the fluid seal and allowing fluid pass from the passageway 408 in the housing 398 of the target connector portion 338 to the passageway 498 in the female connector 332 via the holes 500. When engaged, the resilient seal element 496 exerts a restoring force toward the target connector portion 338 that presses the end of the seal element 496 against the end of the male luer tip 410, forming a fluid seal therebetween. Thus, the fluid can be kept isolated from the external environment while it is transferred from the target connector portion 338 to the female connector 332.

The female connector 332 can be disengaged from the target connector portion 338. The restoring force exerted by the resilient seal element 496 of the female connector 332 causes it to return to its closed position, sealing off its passageway 498. The elastic members 464a, 464b of the resilient member 402 exert a restoring force on the valve member 400, causing the valve member 400 to return to its closed position with its tip 448 abutted against the shelf 416 as the female connector 332 is disengaged.

The '920 Publication discloses additional details and various alternatives that can be applied to the target connector portion 338 of the connector 320.

FIG. 7A shows a perspective view of the syringe 318 and the intermediate connector portion 340 of the connector 320 in an unengaged configuration. FIG. 7B is a top view of the syringe 318 and intermediate connector portion in an engaged configuration. FIG. 7C is a cross-sectional view of the syringe 318 and intermediate connector portion 340 in an engaged configuration. Although FIGS. 7A-7C show the main body 342 of the connector 320 separated from the remainder of the connector 320 for simplicity, it should be understood that the main body 342 can be connected to the remainder of the connector 320 when in use.

In the embodiment shown, the intermediate connector portion 340 is an integral part of the main body 342 of the connector 320. Other configurations are possible. For example, in some embodiments, the intermediate connector portion 340 can be a separate piece connected to the main body 342. The intermediate connector portion 340 can include a female connector 506. In some embodiments, the female connector 506 can have a tapered inner surface. The external surface of the female connector 506 can include external threads 508.

The syringe 318 can have a hollow syringe body 510 defining an internal volume 511. The syringe can include a male luer tip 512 at one end and a shroud 514 surrounding the male luer tip 512. The shroud 514 can have internal threads 516. The male luer tip 512 and threaded shroud 514 can be configured to securely mate with the female connector 506 on the intermediate connector portion 340 of the connector 320, forming a fluid tight connection therebetween. The syringe body 510 can include a body flange 518 positioned at the end of the body opposite the male luer tip 512. The syringe also

includes a plunger 520 that can be slidably received into the internal volume of the syringe body 510. The plunger 522 can include a stopper 522 or other sealing member configured to form a fluid tight seal against the inner surface of the syringe body 510. A plunger flange 524 can be positioned on the plunger 520 at the end opposite the stopper 522.

In some embodiments, the female connector 506 and the male luer tip 512 can be open to the atmosphere when unengaged. Other configurations are possible. For example, in some embodiments, the female connector 506 can be a sealing female connector similar to the female connector 332 described above, and can be for example a version of the Clave® connector. Similarly, the syringe 318 can include a sealing male connector, or a sealing male connector can be connected between the syringe 318 and the female connector 506. In some embodiments the sealing male connector can be a version of the Spiros™ connector. Thus, in some embodiments, the fluid in the syringe 318 and in the connector 320 can be isolated from the environment even when they are disengaged from each other.

In some embodiments, when the syringe 318 is engaged with the connector 320 (as shown in FIG. 7B) the internal volume 511 of the syringe 318 can be in two way fluid communication with the connector 320. Thus, as the plunger 520 is retracted a fluid can be drawn from the connector 320 into the internal volume 511 of the syringe 318. Then as the plunger 520 is advanced the fluid can be directed out of the internal volume 511 and into the connector 320.

As discussed briefly above, the connector 320 can include a source check valve 356 and a target check valve 358. The check valves 356, 358 can function so that when the plunger 520 is retracted the source check valve 356 opens and the target check valve 358 closes, allowing fluid to flow from the vial 314 through the connector 320 and into the syringe 318. Then, when the plunger 520 is advanced the source check valve 356 can close and the target check valve 358 can open, allowing fluid to flow from the syringe 318 through the connector 320 and into the IV bag 316.

FIG. 8A is a perspective view showing the source check valve 356. FIG. 8B is another perspective view showing the source check valve 356 from a different angle. The source check valve 356 can include a disk shaped base 526. A plurality of feet 528 can extend axially from one side of the base 526. In the embodiment shown, the source check valve 356 includes four feet 528, but other numbers of feet 528 can be used, such as three feet, or five feet, or another suitable number of feet 528. In some embodiments, the feet 528 can each be generally cylindrical in shape and can each include a rounded tip 530. Other shapes and configurations are possible. The source check valve 356 can have a sealing surface 531 located on the side opposite the feet 528.

FIG. 9A shows the source connector portion 336, the source check valve 356, and the main body 342 in an exploded cross-sectional view. FIG. 9B is a cross-sectional view of the source connector portion 336, the source check valve 356, and the main body 342 in an assembled configuration with the check valve 356 in an open position. As discussed above, the source connector portion 336 can include a fluid extraction channel 382 having an upper, narrow portion 384 and a lower, wide portion 386. A shoulder 388 can be defined at the transition from the upper portion 384 to the lower portion 386 of the fluid extraction channel 382. In some embodiments, the lower portion 386 of the extraction channel 382 can have a tapered inner surface, so that the lower portion 386 narrows near the shoulder 388. The upper portion 384 can also have a tapered inner surface, so that the upper portion 384 widens near the shoulder 388. In some embodiments, the

21

upper portion 384 and/or the lower portion 386 can be substantially cylindrical or can assume a variety of other shapes having a substantially uniform cross-sectional area.

The main body 342 can include a first fluid passageway 532 leading from the end 534 of the male connector 346 to the end 534 of the intermediate connector portion 340. The first fluid passageway 532 can include an upper portion 536 and a lower portion 538. The lower portion 538 can be wider than the upper portion 536 defining a shoulder 540. The upper portion 536 and lower portion 538 can have tapered or untapered inner surfaces. When assembled, the source check valve 356 can be positioned in the chamber 348 located between the end 534 of the male connector 346 and the shoulder 388 of the fluid extraction channel 382. The source check valve 356 can be positioned so that the feet 528 face toward the end 534 of the male connector 346, while the sealing surface 531 can face toward the shoulder 388. In some configurations, when the pressure in the fluid passageway 332 is sufficiently higher than the pressure in the extraction channel 382, such as when the plunger 520 of the syringe 318 is advanced forcing fluid into the fluid passageway 332, the source check valve 356 is pushed away from the main body 342 and the sealing surface 531 engages the shoulder 388 forming a fluid tight seal that prevents fluid from flowing from the first fluid passageway 532 into the upper portion 384 of the extraction channel 382. In some configurations, when the pressure in the fluid passageway 332 is sufficiently lower than the pressure in the extraction channel 382, such as when the plunger 520 of the syringe 318 is retracted drawing fluid out of the fluid passageway 332, the source check valve 356 is pulled away from the shoulder 388 and the feet 528 rest against the end 534 of the male connector 346 in an open position.

FIG. 10A is a side view of the main body 342 coupled to the source connector portion 336 of the connector 320. FIG. 10B is a cross sectional view of the source connector portion 336 of the connector 320 and the source check valve 356 disposed therein. FIG. 10C is a partial cross-sectional view showing the source check valve 356 positioned in a chamber 348 defined radially by the walls of the female connector 344. FIG. 10D is another cross sectional view showing the source check valve 356 positioned in the chamber 348.

With reference to FIGS. 10A-10D, the base 526 of the source check valve 356 can be disk shaped and can have a diameter d_1 that is less than the diameter d_2 of the chamber 438, defining a space 542 between the side edges of the source check valve 356 and the inner walls of the chamber 348 through which fluid can pass. Also, the feet 528 can be spaced so that open areas 544 are defined between adjacent feet 528.

Thus, when the source check valve 356 is in the open position, fluid can flow from the upper portion 384 of the extraction channel 382, into the chamber 348, through the space 542 between the side edges of the source check valve 356 and the inner walls of the chamber 348, through the open areas 544 between the feet 528, and into the upper portion 536 of the first fluid passageway 532.

In some embodiments, the source check valve 356 can be configured to allow a substantially open flow around the check valve 356 without significant bottlenecking. For example, the space 542 between the side edges of the source check valve 356 and the inner walls of the chamber 348 can have a cross-sectional area A_1 that is at least large as the cross-sectional area A_2 of the upper portion 483 of the extraction channel 382 taken near the chamber 348. This relationship can be expressed as equation (1) below.

$$A_1 \geq A_2$$

(1)

22

In some embodiments, the chamber 348 and the source check valve 356 can both be substantially cylindrical, having diameters d_2 and d_1 respectively (as shown in FIGS. 10C and 10D). The cross sectional area A_1 of the space the space 542 between the side edges of the source check valve 356 and the inner walls of the chamber 348 can then be defined by equation (2) below.

$$A_1 = \pi \left(\frac{d_2}{2} \right)^2 - \pi \left(\frac{d_1}{2} \right)^2$$

In some embodiments, the upper portion 483 of the extraction channel 382 taken near the chamber 348 can be substantially cylindrical and can have a diameter d_3 (as shown in FIG. 10D). The area A_2 can then be defined by equation (3) below.

$$A_2 = \pi \left(\frac{d_3}{2} \right)^2$$

By substituting equations (2) and (3), equation (1) can be rewritten as equation (4) below.

$$\pi \left(\frac{d_2}{2} \right)^2 - \pi \left(\frac{d_1}{2} \right)^2 \geq \pi \left(\frac{d_3}{2} \right)^2$$

By solving equation (4) for d_1 , equation (4) can be rewritten as equation (5) below.

$$d_1 \leq \sqrt{d_2^2 - d_3^2}$$

Thus, when the diameter d_2 of the chamber 348 and the diameter d_3 of the upper portion 483 of the extraction channel 382 are known, the source check valve 356 can be having a diameter that satisfies equation (5) to avoid bottlenecking of fluid as it flows through the space 542.

As shown in FIG. 10D, in some embodiments, when the source check valve 356 is in the open position a space 546 having a height h_1 is defined between the sealing surface 531 and the shoulder 388. The space 546 can allow fluid to flow therethrough. In some embodiments, the source check valve 356 and the chamber 348 can be configured to prevent bottlenecking as fluid flows through the space 546. For example, in the embodiment shown, the smallest area of the space 546 through which the fluid flows can be described as an open an imaginary open cylinder (shown by dotted lines in FIG. 10D) having a height of h_1 , a diameter of d_3 , and a surface area A_3 . If the surface area A_3 of the imaginary cylinder is at least as great as the cross-sectional area A_2 of the upper portion 483 of the extraction channel 382 taken near the chamber 348, bottlenecking can be reduced. This relationship can be expressed as equation (6) below.

$$A_3 \geq A_2$$

The surface area A_3 of the imaginary open cylinder can be expressed as equation (7) below.

$$A_3 = \pi d_3 h_1$$

By substituting equations (3) and (7), equation (6) can be rewritten as equation (8) below.

23

$$\pi d_3 h_1 \geq \pi \left(\frac{d_3}{2} \right)^2 \quad (8)$$

By solving for h_1 , equation (8) can be rewritten as equation (9) below.

$$h_1 \geq \frac{d_3}{4} \quad (9)$$

Thus, when the diameter d_3 of the upper portion **483** of the extraction channel **382** is known, the source check valve **356** can be made to have a total height that is shorter than the height of the chamber **348** by at least

$$\frac{d_3}{4}$$

to reduce bottlenecking as the fluid flows from the upper portion **483** of the extraction channel **382** into the space **546** between the source check valve **356** and the shoulder **388**.

The source check valve **356** can be configured to reduce bottlenecking of the fluid as it flows through the open areas **544** (shown in FIG. 10C) between the feet **528**. For example, if the total area A_4 of the open areas **544** between the feet **528** is at least as great as the cross-sectional area A_2 of the upper portion **483** of the extraction channel **382** taken near the chamber **348**, bottlenecking can be reduced as the fluid flows from the extraction channel **382** and around the check valve **356**. This relationship can be expressed by equation (10) below.

$$A_4 \geq A_2 \quad (10)$$

In the embodiment shown, the feet **528** are arranged so that an imaginary open cylinder (shown by a dotted line in FIG. 10C) can be placed so that its edge intersects each of the feet **528**. The feet **528** can be positioned so that the imaginary cylinder has a diameter d_4 . In some embodiments, the source check valve **356** includes a number n of feet that each have substantially equal diameters d_5 and substantially equal heights h_2 . The area total area A_4 of the open areas **544** can be defined by equation (11) below. It should be noted that because the feet **528** have rounded tips **530**, the area A_4 can be slightly greater than represented by equation (11). In some embodiments, the feet **528** do not have rounded tips and can be substantially cylindrical.

$$A_4 = \pi d_4 h_2 - n d_5 \quad (11)$$

By substituting equations (3) and (11), equation (10) can be rewritten as equation (12) below.

$$\pi d_4 h_2 - n d_5 \geq \pi \left(\frac{d_3}{2} \right)^2 \quad (12)$$

By using feet **528** that satisfy equation (12), bottlenecking can be reduced. For example, if the number n of feet or the diameter d_5 is increased, the height h_2 of the feet can be increased, or the feet can be moved closer to the peripheral edge (increasing d_4) to compensate.

In some embodiments, the source check valve **356** can be configured to provide a substantially uniform flow of fluid. For example, the space **542** between the side edges of the

24

source check valve **356** and the inner walls of the chamber **348** can have a cross-sectional area A_1 that is substantially equal to the cross-sectional area A_2 of the upper portion **483** of the extraction channel **382** taken near the chamber **348**. Similarly, the surface area A_3 of the first imaginary cylinder can be substantially equal to the cross-sectional area A_2 of the upper portion **483** of the extraction channel **382** taken near the chamber **348**. Likewise, the total area A_4 of the open areas **544** between the feet **528** can be substantially equal to the cross-sectional area A_2 of the upper portion **483** of the extraction channel **382** taken near the chamber **348**. The source check valve **356** and chamber **348** can be configured so that other areas of flow also have an area that is substantially equal to the cross-sectional area A_2 of the upper portion **483** of the extraction channel **382** taken near the chamber **348**. For example, in some embodiments, the shoulder **388** or the sealing surface **531** of the check valve **356** can be tapered so that the height of the space **546** is smaller near the side space **542** than near the upper portion **483** of the extraction channel **382**. In some embodiments, the areas discussed herein can be considered to be substantially equal if they vary by an amount less than an acceptable tolerance T . In some embodiments, the acceptable tolerance T can be less than about 1 mm, 0.5 mm, 0.1 mm, 0.05 mm, or 0.01 mm. In some embodiments, the flow areas around the check valve **356** (e.g., A_1 , A_3 , and A_4) can be smaller than A_2 by an amount no larger than tolerance T . Thus, in some embodiments, a small but acceptable amount of bottlenecking can occur as the fluid flows around the source check valve **356**.

In embodiments where the diameter d_2 of the chamber **348** is greater than the diameter d_1 of the source check valve **356**, the source check valve **356** can move not only axially within the chamber, but also radially within the chamber. For example, FIG. 11 shows a cross-sectional view of the source check valve **356** positioned against one side of the chamber **348** when in a closed position. The diameter d_1 of the check valve **356** can be large enough to allow the check valve **356** to adequately seal off the chamber **348** when positioned against one side of the chamber **348**. For example, in some embodiments, the chamber **348** can be generally symmetrical so that the shoulder **388** has a substantially uniform width, and the diameter d_1 of the check valve **356** can be chosen to satisfy the equation (13) below.

$$d_1 > \frac{d_2}{2} + \frac{d_3}{2} \quad (13)$$

In some embodiments, the upper portion **536** of the first fluid passageway **532** can be generally cylindrical in shape and can have a diameter d_6 . In some embodiments, the feet **528** are positioned near enough to the peripheral edges of the check valve **356** so the feet do not drop into the upper portion **536** of the first fluid passageway **532** when the check valve **356** is positioned against the side of the chamber **348**. For example, FIG. 12 shows a cross-sectional view of the source check valve **356** positioned against one side of the chamber **348** in an open position. The feet **528** are positioned so that when the check valve **356** is positioned against one side of the chamber **348** the foot **528a** closest to the first passageway **532** does not drop down into the first passageway **532**. In some embodiments, the feet **528** can be positioned along a circle concentric with the check valve **356**, the circle having a diameter d_4 that satisfies the equation (14) below.

$$d_4 \geq d_6 + d_2 - d_1 \quad (14)$$

25

In some embodiments, the source check valve **356** can have a diameter of about 2 mm to about 20 mm, although diameters outside this range can also be used. A variety of other configurations are possible. For example, the source check valve **356**, the chamber **348**, the extraction channel **382** and/or the first fluid passageway **532** can have non-circular cross sections.

Turning now to FIGS. **13A-B**, FIG. **13A** shows an exploded cross-sectional view of the target connector portion **338**, the target check valve **358**, and the main body **342**. FIG. **13B** shows a cross-sectional view of the target connector portion **338**, the target check valve **358**, and the main body **342** in an assembled configuration with the target check valve **358** in an open position. The main body **342** can include a second fluid passageway **548** that intersects the first fluid passageway **532** at a junction **550**. In some embodiments, the second fluid passageway **548** can intersect the upper portion **536** of the first fluid passageway **532**. In the embodiment shown the second fluid passageway **548** intersects the first fluid passageway **532** at a substantially right angle. Other configurations are also possible. For example, the fluid passageways **532**, **548** can intersect at an oblique angle. The second fluid passageway **548** can have a narrow portion **552** and a wide portion **554** that define a shoulder **556**. In some embodiments, the narrow portion **552** can have a width that is substantially the same as the width of the upper portion **536** of the first fluid passageway **532** near the junction **550**, while in other embodiments the narrow portion **552** can have a width that is smaller or larger than the width of the upper portion **536** of the first fluid passageway **532** near the junction **550**. In some embodiments, the narrow portion **552** and/or the wide portion **554** of the second fluid passageway **548** can have tapered interior surfaces. For example, the wide portion **554** can be tapered so as to receive a tapered male connector **352**, as discussed above.

When assembled, the target check valve **358** can be positioned in the chamber **354** formed between the male connector **352** and the shoulder **556**. In some embodiments, the target check valve **358** can be similar to the source check valve **356** described above, having a disk shaped base **558**, a plurality of feet **560**, and a sealing surface **562**. The target check valve **358** can be positioned with the feet **560** facing the male connector **352** and the sealing surface **562** facing the shoulder **556**. Thus, when the pressure in the second fluid passageway **548** is sufficiently higher than the pressure inside the male connector **352**, such as when the plunger **520** of the syringe **318** is advanced forcing fluid into the main body **342**, the target check valve **358** can be pushed toward the male connector **352** so that the feet **560** rest against the end of the male connector **352** in an open position. When the pressure in the second fluid passageway **548** is sufficiently lower than the pressure inside the male connector **352**, such as when the plunger **520** of the syringe **318** is retracted drawing fluid out of the main body **342**, the target check valve **358** can be pulled away from the main body **342** so that the sealing surface **562** engages the shoulder **556** forming a fluid tight seal that prevents fluid from flowing from the chamber **354** into the narrow portion **552** of the second fluid channel **548**.

In some embodiments, the target check valve **358** and the chamber **354** can be configured to reduce bottlenecks as fluid flows around the target check valve **358** in its open position. For example, the target check valve **358** and chamber **354** can be configured similarly in many ways to the source check valve **358** and chamber **348** described above.

The check valves **356**, **358** can work together to direct fluid through the system. FIG. **14A** shows the flow of fluid (by flow lines) as the plunger **520** is retracted. Fluid is drawn out of the

26

vial **314** through the upper portion **384** of the fluid extraction channel **382**. The fluid flows into the chamber **348** and around the source check valve **356**, which is in the open position. The fluid flows through the first fluid passageway **532** and into the syringe **318**. The fluid can enter the narrow portion **552** of the second fluid passageway **548**, but the target check valve **358** is in the closed position and prevents the fluid from entering the chamber **354**.

FIG. **14B** shows the flow of fluid (by flow lines) as the plunger **520** is advanced. Fluid is expelled from the syringe **318**, into the first fluid passageway **532**, through the narrow portion **552** of the second fluid passageway **548**, into the chamber **354**, around the target check valve **358** (which is in the open position), through the target connector portion **338**, toward the IV bag **316**. The fluid can travel up the first fluid passageway **532** and into the chamber **348**, but the source check valve **356** is in the closed position and prevents the fluid from advancing back into the vial **314**. In some embodiments, the force of the fluid pressing against the source check valve **356** is strong enough to overcome the force of gravity pulling the source check valve **356** downward so as to maintain the source check valve **356** in the closed position.

The check valves **356**, **358** can be formed from rigid, semi-rigid, or deformable materials. In some embodiments, at least the sealing surfaces **531**, **562** of the check valves **356**, **358** can be formed from a material capable of forming a fluid tight seal against a plastic or other rigid material. In some embodiments, the check valves can include a silicon-based deformable material, or a rubber. In some embodiments, the feet **528**, **560** can be formed from different material than the disk shaped base **526**, **558**. In some embodiments, the feet **528**, **560** can be formed from a rigid polycarbonate or other polymeric material.

FIG. **15** is a perspective view of an automated system **600** for transferring fluid, which can be similar to or the same as the other automated fluid transfer systems (e.g., **100**, **200**) disclosed herein. The system **600** can include a base housing **602**, and six transfer stations **604a-f**, located on a front side of the base housing **602**. In some embodiments, the system **600** can include a different number of transfer stations **604a-f** (e.g., one, two, four, five, eight, or more transfer stations). In some embodiments, the transfer stations **604a-f** can be distributed on multiple sides of the base housing **602**. Transfer stations **604b-f** are shown in an empty state having no syringe attached thereto. Transfer station **604a** is shown having a syringe **606** and a connector **608** attached thereto. During operation, a vial (not shown) can be attached to the top of the connector **608** and an IV bag (not shown) can be placed in fluid connection with the connector **608** so that fluid can be transferred from the vial to the syringe **606** and then from the syringe **606** into the IV bag, as discussed above. Also, during operation, some or all of the transfer stations **604a-f** can be equipped similarly to transfer station **604a**. In some embodiments, multiple transfer stations **604a-f** can operate simultaneously. In some embodiments, multiple transfer stations **604a-f** can be placed in fluid communication with a single IV bag so that fluid from multiple vials can be combined into a single IV bag. In some embodiments, one or more of the transfer stations **604a-f** can include a dedicated IV bag so that fluid from only a single transfer stations can be transferred into the dedicated IV bag.

Turning now to FIGS. **16A-16C**, and **17**, a transfer station **604a** is shown in greater detail. FIG. **16A** shows a partial perspective view of the transfer station **604a**, with the syringe **606** and connector **608** in an unengaged configuration. FIG. **16B** shows a left-side view of the transfer station **604a**, with the syringe **606** and connector **608** in an unengaged configuration.

27

ration. FIG. 16C shows a front-side view of the transfer station 604a, with the syringe 606 and connector 608 omitted from view. The transfer station 604a can include an auxiliary housing 610 connected to the base housing 602. The transfer station 604a can also include a top connector piece 612 attached to the base housing 602 above the auxiliary housing 610, and a bottom connector piece 614 attached to the base housing 602 below the auxiliary housing 610. The top connector piece 612 and the bottom connector piece 614 can extend out a distance past the auxiliary housing 610, and a pair of shafts 616a-b can extend vertically between the top connector piece 612 and the bottom connector piece 614. A middle connector piece 618 can be attached to the shafts 616a-b.

The middle connector piece 618 can have a recess 620 configured to receive the syringe body 624. For example, if the syringe body 624 is generally cylindrical, the recess 620 can in the shape of a half cylinder (as shown). The middle connector piece 618 can also include a slit 622 configured to receive the body flange 626 of the syringe 606. The top connector piece 612 can have a recess 628 configured to receive the shroud 630 of the syringe 606 and a portion of the connector 608. In some embodiments, the middle connector piece 618 can be removable, so that it can be interchanged with additional middle pieces (not shown) to provide compatibility with different sizes and shapes of syringes. Also, in some embodiments, the position of the middle connector piece 618 can be adjustable. For example, the middle connector piece 618 can be slid up and down the shafts 616a-b and secured in a variety of location, providing compatibility with syringes of different lengths. In some embodiments, the position of the middle connector piece 618 can be fixed.

The transfer station 604a can include an actuator 632 configured to retract and advance the plunger 634 of the syringe 606. In the embodiment shown, the actuator 632 includes an actuator base 636. Two shafts 648a-b can be positioned at the back of the actuator base 636 and can extend upward from the actuator base 636 into the auxiliary housing 610. Another shaft 640 can be positioned at the front of the actuator base 636 and can extend upward in front of the auxiliary housing 610. An end piece 642 can be attached to the end of the shaft 640 opposite the actuator base 636. The end piece 642 can include a horizontal slit 644 configured to receive the plunger flange 648 of the syringe 606. The end piece 642 can also be configured to receive a portion of the plunger shaft 650 that is near the plunger flange 648. For example, if the plunger shaft 650 includes four longitudinal ribs (as shown), the end piece 642 can include a vertical slit 646 configured to receive one of the longitudinal ribs. The end piece 642 can also include a thumb screw 652 which can be tightened to apply pressure to the plunger flange 648 and prevent the syringe 606 from accidentally disengaging from the transfer station 604a.

In some embodiments, a motor (not shown) is located inside the auxiliary housing 610. The motor can be an electric motor, a pneumatic motor, a hydraulic motor, or other suitable type of motor capable of moving the actuator 632. In some embodiments, the motor can be a piston type motor. In some embodiments, the motor is contained within the base housing 602 rather than in the auxiliary housing 610. In some embodiments, each transfer station 604a-f has an individual motor dedicated to the individual transfer station 604a-f. In some embodiments, one or more of the transfer stations 604a-f share a motor, and in some embodiments, the system 600 includes a single motor used to drive all the transfer stations 604a-f. The motor can drive the shafts 638a-b downward out of the auxiliary housing 610, which in turn drives the rest of the actuator 632 downward causing the plunger 634 to retract

28

from the syringe body 624 to draw fluid into the syringe. The motor can also draw the shafts 638a-b upward into the auxiliary housing 610, which in turn drives the rest of the actuator 632 upward causing the plunger 632 to advance into the syringe body 624 to expel fluid from the syringe.

In some embodiments, the transfer station 604a can include a label 654 that uniquely identifies the specific transfer station 604a. In some embodiments the label 654 can be prominently displayed at the top of the transfer station 604a. The label 654 can be colored, and each of the transfer stations 604a-f can have a different colored label.

The system 600 can include a controller, for controlling the operations of the transfer stations 604a-f. The controller can start and stop the motor(s) of the system 600 to control the amount of fluid that is transferred from the vial to the IV bag at each transfer station 604a-f. The controller can be one or more microprocessors or other suitable type of controller. The controller can be a general purpose computer processor or a special purpose processor specially designed to control the functions of the system 600. The controller can include, or be in communication with, a memory module that includes a software algorithm for controlling the operations of the system 600. The controller can be contained within the base housing 602. In some embodiments, the controller can be external to the base housing 602, and can be for example the processor of a general purpose computer that is in wired or wireless communication with components of the system 600.

In some embodiments, the transfer station 604a includes a sensor (hidden from view in FIGS. 16A-C) configured to determine when the liquid in the vial (not shown) has run out. If the plunger 634 is retracted to draw fluid into the syringe 606 when the vial contains no more fluid, air is drawn out of the vial and travels into the connector 608 toward the syringe. Air may also be drawn into the connector 608 when the vial still contains a small amount of fluid, but the fluid level is low enough that air is drawn out of the vial along with the fluid (e.g., as an air bubble). In some embodiments, the sensor can detect air in the connector 608. For example, the sensor can be an infrared light source (e.g., an LED) and a photodetector, or other form of electric eye.

In some embodiments, the sensor can be located inside the top connector piece 612. The top connector piece 612 can be made from a bottom portion 656 and a top portion 658. FIG. 17 shows a perspective view of the bottom portion 656 of the top connector piece 612, with the top portion 658 removed. The bottom portion 656 can include a central cavity 660 and a pair of grooves 662a-b, one on either side of the recess 628. Grooves 664a-b can connect the grooves 662a-b to the central cavity 660. In some embodiments, the grooves 662a-b, 664a-b can have semi-circular cross sections. In other embodiments, the grooves can be V-grooves, or any other suitably shaped grooves. The grooves 662a-b can be open at the ends furthest from the recess 628. In some embodiments, the grooves 662a-b can also be open at the ends closest to the recess 628. In some embodiments, walls 665a-b can separate the grooves 662a-b from the recess 628, except that the walls 665a-b can have holes 666a-b that connect the grooves 662a-b to the recess 628.

A light source 668 can be located in the groove 662a, and a photodetector 670 can be located in the groove 662b. In some embodiments, the light source 668 can be a laser light source that is aligned to direct a laser beam of light through the hole 666a, across the recess 628, into the hole 666b, and onto the photodetector 670. In some embodiments, the light source 668 can be an LED or other type of light source. In some embodiments, the light source 668, can emit light in many directions, so that some of the light passes through the

29

hole 666a, across the recess 628, into the hole 666b, and onto the photodetector 670. A wire 672 can be connected to the light source 668 and can run along the groove 664a and through the central cavity 660. The wire 672 can provide power or other electric signals from the controller to the light source 668. A wire 674 can be connected to the photodetector 670 and can run along the groove 664b and through the central cavity 660. The wire 674 can carry electric signals from the photodetector 670 to the controller.

In some embodiments, the top portion 658 (not shown in FIG. 17) of the top connector piece 612 can have grooves and/or cavities that correspond to the grooves and/or cavities formed in the bottom portion 656. In some embodiments, the top portion 658 can have a generally flat underside so as to act as a lid to the grooves and/or cavities that are formed in the bottom portion 656. The top portion 658 can be attached to the bottom portion 656 by an adhesive, a clamp, snap or friction fit structures, or various other manners known in the art or yet to be devised. In some embodiments, the top portion 658 is removably attached to the bottom portion 656 so that the user can access the light source 668 and photodetector 670 for calibration, repair, replacement, etc.

When the syringe 606 and connector 608 are attached to the transfer station 604a, the connector 608 (not shown in FIG. 17) can be positioned in the path of light 676 traveling from the light source 668 to the photodetector 670. In some embodiments, the at least a portion of the connector 608 can be made from a substantially transparent plastic or other suitable material that allows the light 676 to pass through the walls of the connector 608. FIG. 18 is a side-view of the syringe 606 and connector 608 and illustrates the location on the connector 608 that intersects the light 676. In some embodiments, the connector 608 can be positioned so that the light 676 passes through the connector 608 at a location that is below the lower end of the source connector portion 677, but above the male luer tip 678 of the syringe 606. This area is marked as region 680 in FIG. 18. In some embodiments, the connector 608 can be positioned so that light 676 passes through the connector 608 above the external shoulder 682 of the connector 608 (shown as region 684). In some embodiments, the connector 608 can be positioned so that light 676 passes through the first fluid passageway 686 at a location above the junction to the second fluid passageway 688 (shown as region 690). In some embodiments, the light 676 passes through the connector 608 near the midpoint between the lower end of the source connector portion 677 and the top of the junction, so that turbulence created as fluid flows in and out of the second fluid passageway 688 does not cause errors in the sensor's readings. In some embodiments, the light 676 passes through the connector 608 at a location that is far enough from the male luer tip 678 of the syringe 606 so that when air is detected as fluid is being drawn into the syringe 606, the flow can be stopped before the air reaches the male luer tip 678.

In some embodiments, the beam of light 676 travelling from the light source 668 to the photodetector 670 is large enough to cover substantially the entire width of the first fluid passageway 686, so that an air bubble cannot travel down into the syringe 606 without crossing the beam of light 676. In some embodiments, the holes 666a-b shown in FIG. 17 can be larger than as shown, or they can be horizontal slits that allow light to intersect substantially the entire width of the first fluid passageway 686.

The light source 668 and photodetector 670 can be configured to detect the presence of air using absorption spectroscopy, emission spectroscopy, scattering spectroscopy, fluorescence spectroscopy, or other suitable manner of

30

distinguishing between the presence of air and the presence of fluid in the path of the beam of light 676.

FIG. 19A is a perspective view of another embodiment of a top connector piece 1900 which can be similar in some regards to the top connector piece 612 described above. FIG. 19B is an exploded view of the top connector piece. The top connector piece 1900 can be used in place of the top connector piece 612 in connection with the automated fluid transfer system 600. For example, the top connector piece 1900 can be connected to the base housing 602 and can function to receive a portion of the syringe 606 or a portion of the connector 608.

The top connector piece 1900 can include a base member 1902 and a cassette 1904. In some embodiments, the base member 1902 can be made of metal, such as aluminum, although other materials can be used. The cassette 1904 can be made from plastic, although other materials can be used. The cassette 1904 can include a bore 1906 configured to align with a bore 1908 formed in the base member 1902 such that the cassette 1904 can be secured to the base member 1902 by inserting a bolt, screw, or other fastener through the bores 1906, 1908. In some embodiments, one or both of the bores 1906, 1908 can be threaded to mate with corresponding threads on the bolt or other fastener. The bore 1906 can include a widened upper portion to receive the head of the bolt therein. The cassette 1904 can also be secured to the base member 1902 by a snap-fit, or friction-fit, or in any other suitable manner.

The base member 1902 can include a cutout region 1910 configured to receive the cassette 1904 such that the top surface of the cassette aligns substantially flush with the top surface of the base member 1902. One or more bores 1912a-c can extend from the back surface of the base member 1902 to the cutout region 1910. In the illustrated embodiment three bores 1912a-c are shown, although it will be understood that other numbers of bores can be used. The outer bores 1912a, 1912c can receive pins or other fasteners used to secure the base member 1902 to the housing 602 of the fluid transfer system 600. The inner bore 1912b can provide a channel that allows wires 1914a-b, 1916a-b to pass from the cutout region 1910 through the base member 1902 and to the housing 602. Many other configurations are possible. For example, a single bore can be used for securing the base member 1902 to the housing 602 and for providing a channel for the wires 1914a-b, 1916a-b.

A first light source 1918a and a corresponding first photodetector 1920a can be positioned inside the top connector piece 1900. The first light source 1918a and first photodetector 1920a can be similar to the light source 668 and photodetector 670 discussed above. Although the first light source 1918a and first photodetector 1920a are located in the cutout region 1910 in FIG. 19B, it will be understood that the first light source 1918a and first photodetector 1920a can be positioned inside of the slots 1922a-b formed in the cassette 1904. The first light source 1918a can be configured to direct light 1924 through a hole 1926a formed in the cassette 1902, across a recess 1928a, through a second hole 1926b formed in the cassette 1902 on the other side of the recess 1928a, and to the first photodetector 1920a. The wire 1914a can provide power or other electric signals from the controller to the first light source 1918a. A wire 1916a can carry electric signals from the first photodetector 1920a to the controller.

The first light source 1918a and first photodetector 1920a can be configured to detect air in the connector 608 similar to the light source 668 and photodetector 670 discussed above. The recess 1928a, 1928b can be configured to receive the syringe 606 and/or connector 608 such that a transparent portion of the connector 608 is positioned in the path of the

31

light **1924** such that the light **1924** passes through a portion of the fluid pathway between the vial and the syringe **606** (e.g., as discussed above in connection with FIG. **18**). The first light source **1918a** and first photodetector **1920a** can be configured to detect air in the fluid pathway and provide a signal to the controller indicating that the vial may need to be replaced.

The portion of the recess **1928a** formed by the cassette can be substantially semicircular in shape to conform to the portion of the connector **608** configured to assign therewith. The portion of the recess **1928b** formed by the base member **1902** can be further enclosed than the portion of the recess **1928a** formed by the cassette, such that a step **1930** is formed on either side of the recess **1928b**. The steps **1930** can facilitate the proper securing and alignment of the connector **608** with the top connector piece **1900**.

A second light source **1918b** and a corresponding second photodetector **1920b** can be positioned inside the top connector piece **1900**. The second light source **1918b** and second photodetector **1920b** can be similar to the light source **668** and photodetector **760** discussed above. Although the second light source **1918b** and second photodetector **1920b** are located in the cutout region **1910** in FIG. **19**, it will be understood that the second light source **1918b** and second photodetector **1920b** can be positioned inside of the slots **1922a-b** formed in the cassette **1904**. The cassette **1904** can have a pair of arms **1934a-b** that extend outwardly, and the slots **1922a-b** can extend along the arms **1934a-b**. The base member **1902** can have corresponding arms **1936a-b** positioned under the arms **1934a-b** of the cassette **1904**. The second light source **1918b** can be configured to direct light **1938** through a hole **1932a** formed in a first arm **1934a** of the cassette **1902**, across a gap formed between the arms **1934a-b**, through a second hole **1932b** formed in the second arm **1934b** of the cassette **1902**, and to the second photodetector **1920b**. The wire **1914b** can provide power or other electric signals from the controller to the second light source **1918b**. A wire **1916b** can carry electric signals from the second photodetector **1920b** to the controller.

In some embodiments, the cassette **1904** can be removable from the base member **1902**, providing access to the light sources **1918a-b**, photodetectors **1920a-b**, and wires **1914a-b**, **1916a-b** for repair or replacement. In some embodiments, the light sources **1918a-b** and/or photodetectors **1920a-b** can be secured to the cassette **1904** and the cassette **1904** can be interchanged with a replacement cassette if a light source **1918a-b** or photodetector **1920a-b** breaks or if different functionality (e.g., a different wavelength of light) is desired.

The second light source **1918b** and the second photodetector **1920b** can be configured to determine whether an IV bag assembly is connected to the connector **608**. In some embodiments, the controller can be configured to abort a command from a user to transfer fluid to an IV bag for a particular transfer station if the controller determines that no IV bag is attached to the particular transfer station, thereby preventing waste of the fluid to be transferred and preventing exposure to potentially hazardous fluids. The controller can also display an error message or alert on the user interface when a command is aborted in this fashion. It should be understood that in some embodiments, a portion of the connector **608** (e.g., target connector portion **338**) can be closed when no IV bag assembly is attached thereto, so that the connector can prevent fluid from escaping when no IV bag assembly is attached. However, if the fluid transfer station is permitted to infuse fluid into the closed connector, high pressure can build up in the connector which can compromise the closed seal of the connector allowing fluid to escape, or can cause damage to the system **600**. The second light source **1918b** and the second

32

photodetector **1920b** are one example of a sensor configured to determine whether an IV bag assembly is attached to the connector **608**, and it will be understood that other sensor types (e.g., weight sensors) can also be used for detecting the presence of the IV bag assembly.

The manner in which the second light source **1918b** and the second photodetector **1920b** detect the presence of an IV bag assembly will be described in connection with FIGS. **19C-E**. FIG. **19C** is a side view of a connector **1950** which can be similar to the connector **320** or any other connector described herein. The connector **1950** can include a source connector portion **1952** and a target connector portion **1954**. In the illustrated embodiment, the source connector portion **1952** and the target connector portion **1954** can be attached to a main body piece **1956** which can have an intermediate connector portion **1958** configured to receive a syringe or other intermediate measuring container.

FIG. **19D** is a cross sectional view of the connector **1950** that shows the target connector portion **1954** in a closed state. FIG. **19E** is a cross sectional view of the connector **1950** that shows the target connector portion **1954** in an open state. The target connector portion **1954** can be similar to the target connector portion **338** described herein. The target connector portion **1954** can include a housing **1960** and an end cap **1962** that includes an elongate plunger **1964**. A valve member **1966** can be slidably engaged with the plunger **1964** such that when the valve is in the closed position, as shown in FIG. **19D**, the base **1968** of the valve member **1966** overlaps only the end of the plunger **1964**, leaving at least a portion of the plunger **1964** exposed. When the connector **1965** of the IV bag assembly is attached to the target connector portion **1954** (e.g., as described in connection with FIGS. **6D-E**), the valve member **1966** is displaced toward the end cap **1962** as shown in FIG. **19E**.

The second light source **1918b** and the second photodetector **1920b** are shown schematically in FIGS. **19D-E**. In some embodiments, at least a portion of the housing **1960** and at least a portion of the plunger **1964** can be made of a material that is transparent to the light **1938** emitted by the second light source **1918b**, while the valve member **1966** can be made of a material that is opaque to the light **1938**, or otherwise prevents the light **1938** from reaching the second photodetector **1920b** when placed in the path of the light **1938**. Thus, when no IV bag assembly is attached to the connector **1950** and the target connector portion **1954** is in the closed configuration (as shown in FIG. **19D**), the light **1938** can pass through the transparent housing **1960**, through the transparent plunger **1964**, and to the second photodetector **1920b**. When the second photodetector **1920b** detects the light **1938** it can send a signal to the system controller indicating that no IV bag assembly is attached to the target connector portion **1954**. When the connector of an IV bag assembly is attached to the target connector portion **1954** the base **1958** of the valve member **1966** can intersect the path of the light **1938** and prevent the light **1938** from reaching the second photodetector **1920b**, as shown in FIG. **19E**. When the second photodetector **1920b** does not detect light **1938**, it can send a signal to the system controller indicating that target connector portion **1954** is in the open configuration and an IV bag assembly is attached thereto.

In some embodiments, the connector **1950** can be aligned so that the light **1938** passes through the open space **1970** next to the plunger **1964** without intersecting the plunger **1964**. Thus, in some embodiments, the plunger **1964** can be made of a material that not transparent to the light **1938**. In the open configuration, as shown in FIG. **19E**, the base **1968** of the valve member **1966** fills the space **1970** adjacent to the

33

plunger 1964 to block the light 1938. Thus, in some embodiments, the light 1938 does not pass through the fluid flow path 1972 formed through the target connector portion 1954, which can be advantageous in certain circumstances such as when a fluid is transported through the connector that would prevent the light 1938 from reaching the second photodetector 1920b.

FIGS. 19D-E also illustrate the light 1924 emitted by the first light source 1918a being transmitted through the fluid flow path 1974 formed between the vial and the syringe to the first photodetector 1920a, as described above.

Returning now to FIG. 15, the system 600 can include a user interface 692 for receiving information and commands from the user and for providing information to the user. The user interface 692 can be part of an external unit 694, or it can be integrated into or attached to the base housing 602. The user interface 692 can include, for example, a touch screen display. The user interface 692 can be in wired or wireless communication with the controller. In some embodiments, a cable 696 connects the external unit 694 to the base housing 602 and provides a communication link between the user interface 692 and the controller. In some embodiments, the controller can be contained in the external unit 694 along with the user interface 692 and the controller can send and receive signals to and from components (e.g., the motors) of the system 600 through the cable 696. The user interface 692 can be configured to receive instructions from the user regarding the amounts of fluids to be transferred by the transfer stations 604a-604f. The user interface 692 can deliver the instructions to the controller to be stored in a memory and/or used to actuate the motor(s) to transfer the desired amount of fluids.

In some embodiments, the system 600 can include a communication interface (shown schematically in FIG. 15 as antenna 691). The communication interface 691 can be configured to provide a communication link between the controller and a remote source, such as a remote terminal or an automated management system. The communication link can be provided by a wireless signal or a cable or combination of the two. The communication link can make use of a network such as a WAN, LAN, or the interne. In some embodiments, the communication interface can be configured to receive input (e.g., fluid transfer commands) from the remote source and can provide information (e.g., results or alerts) from the controller to the remote source. In some embodiments, the remote source can be an automated management system which can coordinate actions between multiple automated fluid transfer systems (e.g., 100, 200, and 600).

The system 600 can also include a bar code scanner 698, in communication with the controller and/or memory. The bar code scanner 698 can be used to provide information about the system 600 to the controller and/or the memory. For example, the syringe 606 can include a bar code that identifies the size and type of the syringe 606. The user can scan the syringe 606 with the bar code scanner 698 and then scan a bar code associated with the transfer station 604a to inform the controller of the size of the syringe 606 that is attached to the transfer station 604a. Different sizes of syringes can hold different volumes of fluid when their plungers are withdrawn by the same distance. Thus, when the controller is tasked with filling the syringe 606 with a predetermined amount of fluid, the controller can determine how far the plunger is to be withdrawn to fill the particular type of syringe with the predetermined amount of fluid. The vials (not shown) can also include bar codes that indicate the type of fluid contained therein. The user can scan a vial and then scan the bar code associated with the particular transfer station the vial is to be installed onto. Thus, the controller can be aware of what fluids

34

are controlled by which transfer stations to facilitate automated transfer of fluids. Other components of the system 600 can also include bar codes readable by the bar code scanner 698 for providing information about the components to the controller and/or memory. In some embodiments, the user interface 692 can be configured to allow the user to input data relating to the size of the syringe 606, the type of fluid contained in a vial, etc. instead of using the bar code scanner 698.

FIG. 20 is a perspective view that schematically shows another embodiment of an automated fluid transfer system 2000. Some aspects of the automated fluid transfer system 2000 can be similar to or the same as the other automated fluid transfer systems (e.g., 100, 200, and 600) described above. The automated fluid transfer system 600 can include a base housing 2002, and six transfer stations 2004a-f (although the system 600 can have other numbers of transfer stations). In FIG. 20, the transfer stations 2004a-f are shown schematically as boxes, but it should be understood that each of the transfer stations 2004a-f can include structure similar to or the same as that described above in connection with the transfer station 604a. For example, each transfer station can include a fluid transfer subsystem (e.g., subsystem 300 or 1900) including a vial, a syringe, and an IV bag assembly.

The automated fluid transfer system 2000 can include a support bar assembly 2050. FIG. 21 is a side view schematically showing a portion of the support bar assembly. With reference now to FIGS. 20 and 21, the support bar assembly 2050 can include a substantially horizontal support bar 2052, supported on either side by an arm 2054. Each arm 2054 can be attached to the side of the base housing 2002 by an attachment piece 2056. In some embodiments the attachment piece can be integrally formed with the base housing 2002 or secured thereto, for example, by an adhesive or by one or more screws 2055 or other fasteners. The arm 2054 can be attached to the attachment piece 2056 by a shoulder bolt 2058, so that the arm 2054 can pivot on the shoulder bolt 2058. The rotational range of the arm 2054 can be limited by an upper dowel pin 2060 and a lower dowel pin 2062. A spring plunger 2064 can be positioned on the arm 2054 and can be configured to slide into one or more locking holes (hidden from view in FIGS. 20 and 21) to lock the arm 2054, and the support bar 2052, in position. The spring plunger 1064 can be pulled out of the locking hole to release the arm 2054 from the locked position. In FIGS. 20 and 21, the arms 2054 and support bar 2052 are shown locked in an upward position with the arm 2054 positioned adjacent to the upper dowel 2060. The support bar 2052 can be configured to hold or otherwise support at least a portion of the one or more fluid transfer subsystems of the fluid transfer stations 2004a-f. For example, when locked in the upward position, the support bar 2052 can be positioned so that the target connector portion, the female connector attached to the target connector portion, the IV bag, or other portion of the IV bag assembly can rest on the support bar 2052 to reduce the amount of stress placed on the connector.

FIG. 22 is a partial perspective view that schematically shows another embodiment of an automated fluid transfer system 2200 that, in some regards, can be the same as or similar to the other automated fluid transfer systems (e.g., 100, 200, 600, and 2000) disclosed herein. In some embodiments, one or more of the transfer stations (e.g., 2204a) can include a support arm 2250. The support arm 2250 can be integrally formed with or attached to the top connector piece 2212. Alternatively, the support arm 2250 can be separate from the top connector piece 2212 and can be secured, for example, directly to the base housing 2202 by one or more screws or other fasteners. In some embodiments, the support

35

arm **2250** can be substantially “L” shaped, having an elongate extension portion **2252** and a support platform **2254**. The support platform **2254** can be configured to hold or otherwise support at least a portion of the fluid transfer subsystems of the fluid transfer station **2204a**. For example, the support platform **2254** can be positioned so that the target connector portion **2236**, the female connector (not shown in FIG. **22**) attached to the target connector portion **2236**, the IV bag (not shown in FIG. **22**), or other portion of the IV bag assembly can rest on the support platform **2254** to reduce the amount of stress placed on the connector.

In some embodiments, the support arm **2250** can include a weight sensor **2256**, or other type of sensor, capable of determining whether an IV bag assembly (not shown in FIG. **22**) is connected to the target connector portion **2236**. For example, the weight sensor **2256** can “feel” the weight of the IV bag as the support arm **2250** provides support thereto. The weight sensor **2256** can be in electronic communication with the controller so that the controller can confirm that an IV bag assembly is attached to the target connector portion **2236** before transferring fluid into the IV bag.

In some embodiments, the weight sensor **2256** can be used to confirm that the correct amount of fluid was transferred to the IV bag. The controller can be configured to calculate an expected weight for the IV bag from the instructions received from the user and from information stored in a memory, e.g., the amount of fluid to be transferred, the density of the fluid to be transferred, the starting weight of the empty IV bag, etc. Once the transfer of fluid is complete the controller can measure the final weight of the IV bag using the weight sensor and can compare the final weight to the expected weight. If the final weight differs from the expected weight by more than an acceptable tolerance amount (e.g., determined by the accuracy of the weight sensor), the controller can send an error message or alert to the user interface informing the user that an error likely occurred in the fluid transfer (e.g., the wrong fluid type was transferred or the wrong amount of fluid was transferred).

FIG. **22A** is a partial perspective view of another embodiment of an automated fluid transfer system **2270** that, in some regards, can be the same as or similar to the other automated fluid transfer systems (e.g., **100**, **200**, **600**, **2000**, and **2200**) disclosed herein. The system **2270** can include a tray **2272** extending out from the housing **2274**. The tray **2272** can be configured to support the IV bag **2276**. The tray **2272** can have flat base **2278** and sides **2280a-b** that turn up (e.g., by about 30° to about 60°) to prevent the IV bag **2276** from sliding off the side of the tray **2272**. The end **2282** of the tray **2272** furthest from the housing **2274** can be open, having no turned up side, so that the IV bag can hang over the edge of the tray **2272**. A support foot **2279** can extend from the base of the housing **2274** to prevent the system **2270** from tipping forward under the weight of the IV bag **2276**.

The tray **2272** can include a hole or cutout **2284** configured to align with the target connector portion **2286** of the connector (which can be similar to the connector **320** or any other connector disclosed herein). In some embodiments, the outer housing **2288** of the target connector portion **2286** can rotate relative to the connector **2290** (which can be similar to the female connector **322**) of the IV bag assembly. Because at least a portion of the target connector portion **2286** is rotatable, the connector **2290** is not required to rotate when it is attached or detached to the target connector portion **2286**, so that the tubing **2292** is not twisted or kinked and the IV bag **2276** need not be twisted. In some embodiments, the target connector portion **2286** can rotate to engage the connector **2290** in a manner similar to that described above in connec-

36

tion with FIGS. **6D-E**, although it will be understood that any rotating connector can be used. The hole or cutout **2284** formed in the tray **2272** can be configured to allow a user's hand to pass through therethrough when rotating the housing **2288** of the target connector portion **2286**.

The tray **2272** can be removably secured to the housing **2274**. In some embodiments, the tray **2272** can be bolted, screwed, or otherwise fastened to the housing **2274**. A snap fit connection or a friction-fit connection can also be used. In some embodiments, the end of the tray can fit between the top connector piece **2294** and the auxiliary housing **2296** of the transfer station with which the tray **2272** is associated. The embodiment illustrated in FIG. **22A** shows a single tray **2272** attached to a transfer station of the system **2270**, but it will be understood that a plurality of individual trays can be used, each tray being associated with one of the transfer stations. In some embodiments, a single tray can be used for more than one or all the transfer stations.

FIG. **23** is a flowchart that schematically shows a method **2300** of operation for an automated fluid transfer system (e.g., **100**, **200**, **600**, **2000**, and **2200**). At block **2302**, the system receives a fluid transfer command. The fluid transfer command can be received, for example, via a user interface from inputs provided by a user, or via a communication interface from a remote terminal or an automated management system. The fluid transfer command can include information such as a fluid type to be transferred, an amount of the fluid to be transferred, and a desired concentration of the fluid. In some embodiments, a fluid transfer command can include information for multiple fluids to be combined into a compounded mixture.

At block **2304**, the controller determines whether the fluid transfer stations of the system are currently equipped to transfer the requested fluids. In some embodiments, the system includes a memory that includes, for example, a database or lookup table so that the controller can determine the type of fluids associated with each transfer station. If the fluid transfer stations do not have the specified fluid, the method can proceed to block **2306** wherein the user interface can prompt the user to change the fluid(s) of the fluid transfer station(s). In some embodiments, the controller can determine a recommended fluid to replace (e.g., using a history of usage stored in the memory) and provide the recommendation to the user via the user interface. After the user makes the changes to the fluid transfer station(s), the method **2300** can return to block **2304** to confirm that the transfer station(s) are properly equipped.

In some embodiments, the user can specify one or more transfer stations to use for the fluid transfer, rather than specifying the types of fluids desired. Thus in some embodiments, blocks **2304** and **2306** can be omitted. In some embodiments, the user interface can display to the user the types of fluids associated with the different transfer stations to aid the user in selecting the transfer stations to use for the fluid transfer.

In some embodiments, the system can contain concentrated fluids in the source containers and in some circumstances the fluids are to be diluted with a diluent prior to delivery to the patient. Therefore, in some instances, the controller can determine a desired amount of diluent based upon the concentration of the fluid in the source container, the desired concentration, and the amount of fluid to be transferred. The user interface can prompt the user to fill the target IV bag with the desired amount diluent. Alternatively one or more of the transfer stations of the system can include diluents. Thus, in some embodiments, the controller will determine whether transfer stations are equipped with the desired medication and the desired diluent.

37

If the fluid transfer stations are properly equipped, the method **2300** can proceed to block **2308** where the controller determines whether the IV bag assembly is properly attached. In some embodiments, the system can include, for example, a weight sensor or IR sensor capable of determining whether the target connector portion for a transfer station is connected to an IV bag assembly. In some embodiments, the weight sensor and controller can determine whether the IV has been filled with a desired amount of diluent. In some embodiments, the memory can include a database or lookup table indicating which transfer stations are associated with which IV bags (which can be especially useful when multiple transfer stations are associated with a single IV bag). The information can be input by the user via the user interface or by scanning bar codes on the IV bags and transfer stations. If the controller determines that the IV bag assembly is not properly attached (e.g., no IV bag attached, or incorrect IV bag weight for desired diluent, or a wrong combination of transfer stations associated with the IV bag), the user interface can prompt the user to attach an IV bag or otherwise change the IV bag configuration. After the user makes the changes, the process **2300** can return to block **2308** to confirm that the IV bag assembly is properly attached and configured.

If the IV bag assembly is properly attached, the process **2300** proceeds to block **2312** where the system transfers fluid(s) from the transfer station(s) to the IV bag, as will be described in greater detail below.

FIG. **24** is a flowchart that schematically shows an embodiment of a method **2400** for transferring an amount of fluid from a vial to an IV bag. At block **2402**, the controller determines an amount of fluid to be transferred. In some embodiments, the amount can be specified directly by the fluid transfer command. In some embodiments, the amount of fluid (e.g., medication or diluent) can be affected by the desired concentration and the concentration of the fluid contained in the vial.

At block **2404**, the controller determines whether the transfer amount is greater than the effective maximum volume of the syringe associated with the transfer station. In some embodiments, the memory can include a database or lookup table that stores the sizes of the syringes associated with the different transfer stations. The information can be input by the user via the user interface or by scanning bar codes on the syringes and transfer stations. In some embodiments, the effective maximum volume of a syringe is the volume of the syringe when the plunger is substantially fully retracted. In some embodiments, the effective maximum volume of the syringe is the volume of the syringe when the plunger is retracted by the maximum amount that the actuator is able to retract.

If the amount to be transferred is greater than the effective maximum volume of the syringe, the method **2400** proceeds to block **2406** where the controller causes the plunger of the syringe to be withdrawn so as to draw the effective maximum volume of fluid from the vial into the syringe. As the fluid is transferred to the syringe in block **2406**, the system can monitor for air bubbles, in block **2408**, which can indicate that the fluid in the vial has run out. If a bubble is detected at block **2408**, the method **2400** can interrupt block **2406** and prompt the user to replace the empty vial at block **2410**. Once the vial has been replaced, the method **2400** can return to block **2406** and finish filling the syringe.

Once the syringe has been filled the method can proceed to block **2412** where the system determines whether an IV bag is attached to the target connector portion of the relevant transfer station. In some embodiments, a weight or IR sensor can be used to detect the presence of an IV bag or a connector

38

attached to the target connector portion. Because an IV bag can be disconnected by mistake during a fluid transfer, in some embodiments the system can be configured to check for a connected IV bag each time the plunger of the syringe is to be advanced to drive fluid out of the syringe. In some embodiments, the system checks for an attached IV bag only at the start of the fluid transfer, so blocks **2412** and **2414** can be omitted. If the IV bag is not attached, the method **2400** can proceed to block **2414** where the user interface can prompt the user to reattach the IV bag. In some embodiments, the UI can provide an alert message to the user indicating that an error has likely occurred (e.g., an IV bag was removed prematurely). Once the changes have been made, the method **2400** can return to block **2412** to confirm that the IV bag is properly attached. In some embodiments, if the IV bag is not properly attached, the method **2400** can abort the fluid transfer, rather than proceeding to block **2414**, and display an error message or alert to the user.

Once the system determines that the IV bag is attached, the method **2400** can advance to block **2416** where the controller can cause the actuator to advance the plunger of the syringe to drive the fluid out of the syringe and into the IV bag. At block **2418**, the method can subtract the effective max volume of the syringe (i.e., the amount added to the IV bag at block **2416**) from the amount of fluid to be transferred. Then the method **2400** can return to block **2404**.

If, at block **2404**, the controller determines that the amount to be transferred is less than the effective maximum volume of the syringe, the method **2400** can advance to block **2420** where the controller causes the actuator to withdraw the plunger of the syringe by a distance to draw the remaining transfer amount of fluid into the syringe. The controller can be configured to determine the distance to draw back the plunger based on the amount fluid remaining to be transferred and by the size of the syringe, which can be stored in a database or lookup table in the memory.

At block **2422**, the system can monitor for air bubbles similarly to block **2408**. If an air bubble is detected, the process **2400** can interrupt block **2420** and proceed to block **2424** where the user interface can prompt the user to replace the empty vial. Once the vial has been replaced the method **2400** can return to block **2420** and finish filling the syringe with the desired amount of fluid.

Once the syringe contains the remaining fluid to be transferred, the process can advance to block **2426**, where the system determines whether an IV bag is attached similar to block **2412**. If no IV bag is properly attached, the method **2400** can advance to block **2428**, where the user interface can prompt the user to reattach the IV bag. Once the changes have been made the method **2400** can return to block **2426** to confirm that an IV bag is properly attached. Then the method **2400** can advance to block **2430** where the controller can cause the actuator to advance the plunger of the syringe to drive the fluid from the syringe into the IV bag.

The method **2400** can end at block **2432**. In some embodiments, the method **2400** can repeat for one or more additional fluids (e.g., a diluent or additional medication for a compounding procedure) transferred from one or more additional transfer stations. In addition, the blocks and order illustrated are exemplary methods. Modification is also possible. For example, the system can detect whether a bag is attached (e.g., blocks **2412**, **2426**) prior to drawing fluid into the syringe (e.g., blocks **2406**, **2420**).

FIG. **25** is a flowchart that schematically shows an embodiment of a method **2500** for confirming the successful transfer of fluid by checking the weight of the final IV bag. At block **2502**, the controller can determine an expected IV bag weight

for the final IV bag filled with the transferred fluid. The expected weight can be determined by the starting weight of the empty IV bag (or the starting weight of the IV bag with diluent), and the amount and density of fluid to be transferred into the IV bag.

At block **2504**, the system can measure the actual IV bag weight. In some embodiments, the system can include a weight sensor and can automatically measure the weight of the IV bag once the fluid transfer is complete. In some embodiments, the user interface can prompt the user to weigh the IV bag and enter the weight. In some embodiments, the user interface can prompt the user that the transfer is complete and display the expected weight for the IV bag. The user can then weigh the IV bag and compare the actual weight against the displayed expected weight.

At block **2506**, the controller can compare the actual IV bag weight to the expected IV bag weight. If the actual IV bag weight differs from the expected IV bag weight by more than a threshold tolerance amount, the method **2500** can determine that an error occurred during the fluid transfer and advance to block **2510**. At block **2510**, the controller can attempt to determine possible causes of the fluid transfer failure. Many circumstances can lead to a fluid transfer failure. For example, if the user changes the type of fluid for a fluid transfer station without properly updating the database, the IV bag can contain the correct amount of fluid but since the fluid can have a different density the final weight of the IV bag can be different from the expected amount. If the user changes the syringe size for the transfer station without properly updating the database the actuation of the plunger can transfer an amount of fluid different than intended and the final weight of the IV bag can differ from the expected weight. The controller can be configured determine possible causes for the failure based at least in part on the amount by which the actual IV bag weight differs from the expected weight. At block **2512**, the user interface can inform the user of the failure and can display one or more possible causes for the failure to aid the user in trouble shooting the problem.

If the actual IV bag weight is within the threshold tolerance amount of the expected weight, the system can conclude that the fluid was transferred successfully, and the method can advance to block **2508**. At block **2508**, the user interface can inform the user that the fluid was transferred successfully. The threshold tolerance amount can be determined by several factors, including the precision of the weight sensors, the amount of fluid transferred, and the accuracy provided by the syringe(s) used. It should be noted that some fluid transfer errors can go undetected by checking the weight of the IV bag. For example, if an incorrect fluid is used that has the same density as the correct fluid, the final IV bag will weigh the correct amount. However, by checking the weight of the IV bag, many errors can be detected.

FIG. **26** is a partial sectional view that schematically shows another embodiment of a fluid transfer subsystem **2600** that can include a vial **2614**, a syringe **2618**, and a connector **2620**. In some embodiments, the vial **2614**, syringe **2618**, and connector **2620** shown in FIG. **26** can be the same as or similar to, for example, to the vial **314**, syringe **318**, and connector **320** described above. In some embodiments, the connector **2620** can include a main body portion **2642**, a source connector portion **2636** configured to connect to the vial **2614**, a target connector portion **2638** (partially shown in FIG. **26**) configured to connect to an IV bag assembly (not shown in FIG. **26**), and an intermediate connector portion **2640** configured to connect to the syringe **2618**.

In some embodiments, the source connector portion **2636** can similar to the source connector portion **336** described

above. The source connector portion **2636** can be integrally formed with the main body portion **2642** of the connector **2620**, or the source connector portion **2636** can be separately formed and secured to the main body portion **2642**, for example, by a plastic welding adhesive or other manner as described above. In some embodiments, the source connector portion **2636** includes a piercing member **2670** which can include an elongate shaft **2672** and pointed tip **2674**. The piercing member **2670** can be configured to puncture a septum **2660** formed in a cap **2659** of the vial **2614** when the vial **2614** is pressed onto the connector **2620**.

In some embodiments, the source connector portion can include a fluid extraction channel **2682** extending from an extraction aperture **2683** formed in a portion of the piercing member **2670** to the main body portion **2642** of the connector **2620**. The fluid extraction channel **2682** can be configured to allow fluid **2666** to flow out of the vial **2614** and into the connector **2620**, e.g., when the plunger **2619** of the syringe **2618** is withdrawn. In some embodiments, the connector **2620** can include a source check valve **2656** formed therein and configured to allow fluid to flow from the vial into the connector **2620** and prevent fluid from flowing from the connector **2620** into the vial **2614**. In some embodiments, the source check valve **2656** can be similar to the check valve **356** described above or it can be a duckbill valve formed in the fluid extraction channel **2682**, as schematically shown in FIG. **26**. Many other variations are possible.

The source connector portion **2636** can also include a regulator channel **2690** extending from a regulator aperture **2692** up through a portion of the elongate shaft **2672** to an opening **2693** formed in the piercing member **2670**. The regulator channel **2690** can allow air to enter the connector **2620** and flow into the vial **2614** as the fluid **2666** is withdrawn, thereby maintaining a substantially constant pressure inside the vial **2614**. In some embodiments, a regulator check valve **2655** can be formed in the regulator channel **2690** to prevent fluid **2666** from escaping from the vial **2614** via the regulator channel **2690**. The connector **2620** can also include a filter **2661** formed over the regulator aperture **2692** to prevent contaminants or other foreign particles from entering the regulator channel **2690** and contacting the fluid **2666**. In some embodiments, the filter **2661** can be permeable to air so that air is permitted to enter the vial **2614** via the regulator channel **2690**. In some embodiments, the filter **2661** can be impermeable to the fluid **2666** and can be used in conjunction with, or in place of, the regulator check valve **2655** to prevent fluid **2666** from exiting the vial **2614** via the regulator channel **2690**.

In some embodiments, the source connector portion **2636** can differ from the source connector portion **336** by not including a bag to hold the air that enters the vial **2614**. Thus, the air that enters the vial **2614** can directly contact the fluid **2666** contained therein. In some embodiments, the connector portion **2636** is only used for vials **2614** containing fluid **2666** that will not react with, or otherwise be adversely affected by, the air. In some embodiments, the filter **2661** and/or regulator check valve **2655** can be configured to allow only certain gases, which will not adversely affect the fluid **2666**, to enter the vial **2614**.

The target connector portion **2638** can be similar to the target connector portion **338** described above, the disclosure of which applies to the embodiment shown in FIG. **26**. Only the main connector portion **2652** of the target connector portion **2638** is shown in FIG. **26**. The target connector portion can be configured to provide fluid communication between the connector **2620** and an IV bag assembly (not shown in FIG. **26**) similar or the same as the IV bag assembly **330**

41

described above. The connector 2620 can include a target check valve 2658 configured to allow fluid to flow from the connector into the IV bag assembly, e.g., when the plunger 2619 of the syringe 2618 is advanced, and prevent fluid from flowing from the VI bag assembly into the connector 2620. The target check valve 2658 can be similar or the same as the target check valve 358 described above, or it can be a duckbill valve as shown schematically in FIG. 26.

The intermediate connector portion 2640 can be configured to removably receive the syringe 2618 and provide a sealed fluid pathway between the connector 2620 and the syringe 2618. In some embodiments, the intermediate connector portion 2640 can be the same as or similar to the intermediate connector portion 340 described above.

The fluid transfer subsystem 2600 can be used as a fluid transfer station on an automated fluid transfer system, which can be, for example, similar to the automated fluid transfer system 600 described above.

FIG. 27A is a perspective view of an embodiment of a fluid transfer module in the form of a connector 2700, which can be similar in many regards to the connector 320 or any other connector disclosed herein. FIG. 27B is another perspective view of the connector 2700. The connector 2700 can be used to transfer fluid from a source container (e.g., a vial) to an intermediate measuring container (e.g., a syringe) and then to a target container (e.g., an IV bag). The connector 2700 can include a source connector portion 2702 configured to interface with the source container (e.g., a vial), an intermediate connector portion 2704 configured to interface with the intermediate measuring container (e.g., a syringe), and a target connector portion 2706 configured to interface with the target container (e.g., an IV bag assembly).

The connector 2700 can function to transfer fluid from the source container to the target container similarly to the connector 320 or the connector 2600 or any other connector disclosed herein. Fluid can be extracted from a vial (not shown) through the fluid extraction aperture 2708, and air can enter the vial via the air inlet 2710 and air outlet 2712 to replace the volume of extracted fluid. The fluid extracted from the vial can be drawn through the connector 2700 and into the syringe (not shown) via the opening 2714 formed in the intermediate connector portion 2704. A source check valve (hidden from view in FIGS. 27A-B) can be configured to allow fluid to flow from the fluid extraction aperture 2708 to the opening 2714 in the intermediate connector portion 2704 while preventing fluid from flowing in the reverse direction back into the vial. The fluid can be driven from the syringe into the connector 2700 via the opening 2714, and the fluid can be directed into the target connector portion 2706 and into an IV bag assembly (not shown) attached to the target connector portion 2706. A target check valve (hidden from view in FIGS. 27A-B) can be configured to allow the fluid to flow from the opening 2714 in the intermediate connector portion 2704 to the target connector portion 2706 while preventing fluid from flowing in the reverse direction.

FIG. 28A is an exploded perspective view of the connector 2700. FIG. 28B is another exploded perspective view of the connector 2700. The connector 2700 can include an upper housing member 2720 and a lower housing member 2722. The upper housing member 2720 can include the source connector portion 2702 of the connector 2700, and the lower housing member 2722 can include the intermediate connector portion 2704 of the connector 2700.

The upper housing member 2720 can include a piercing member 2724 made up of an elongate substantially cylindrical shaft 2726 and a pointed tip 2728. The piercing member 2724 can be configured to pierce the septum of a vial (not

42

shown) when the vial is attached thereto. The upper housing member 2720 can include retaining arms 2730a-b configured to secure the vial to the connector 2700, as described herein. The piercing member 2724 can include a fluid extraction aperture 2708 formed on one side thereof. The fluid extraction aperture can be a slit that extends from near the end of the pointed tip 2728 down onto the shaft 2726, although openings of other shapes can also be used. In some embodiments, the slit shape can facilitate the full extraction of fluid from the vial. A fluid pathway 2732 can extend from the fluid extraction aperture 2708 to a fluid outlet opening 2734 formed in the bottom surface of the base 2736 of the upper housing member 2720. The piercing member 2724 can also include an air outlet 2712 that allows air to enter the vial as fluid is extracted therefrom to equalize the pressure differential caused by the extraction of fluid. The air outlet 2712 can receive air from an air pathway 2738 that extends through the shaft 2726 and through the base 2736 and to an air inlet opening 2740 formed in the base 2736 of the upper housing 2720.

The upper housing member 2720 can include a female end 2742 configured to receive a male end 2744 of the target connector portion 2706. The target connector portion 2706 can be similar to the other target connector portions described herein (e.g., 338), the disclosure of which applies also to the target connector portion 2706. The male end 2744 can be secured to the female end 2742 by applying a plastic welding adhesive (such as Dichloromethane) to the outer surface of the male end 2744 and/or to the inner surface of the female end 2742 before insertion. The Dichloromethane can chemically weld the outer surface of the male end 2744 to the inner surface of the female end 2742. Other methods can be used to connect the male end 2744 to the female end 2742, such as sonic welding, threading, adhesives, etc. It will also be understood that the target connector portion can include the female end of the interface while the top housing member can include the male end thereof. Indeed, any suitable interface for securing the target connector portion 2706 to the upper housing member 2702 can be used. In some embodiments, the connection between the male end 2744 and the female end 2742 is hermetically sealed, and in some embodiments includes a sealing member (not shown), such as an O-ring, to provide the hermetic seal. A fluid pathway 2746 can extend from the opening in the female end 2742 to a fluid inlet opening 2748 formed in the bottom surface of the base 2736 of the upper housing member 2720.

The lower housing member 2722 can include a chamber 2750 enclosed by a base wall 2752 and by side walls 2754 having an open top. The chamber 2750 can be configured to receive the base 2736 of the upper housing member 2720 when the top housing member 2720 is secured to the bottom housing member 2722. The side walls 2754 can include projections 2756a-b formed near the top thereof, which can be configured to mate with corresponding slots 2758a-b formed in the upper portion of the base 2736 for provide a snap-fit connection between the top housing member 2720 and the bottom housing member 2722. It will be understood that the top housing member 2720 can be secured to the bottom housing member 2722 using various other techniques including an adhesive, sonic welding, a friction-fit, or any other suitable manner. The side walls 2754 of the lower housing member 2722 can include a front cutout 2760 configured to receive a portion of the female end 2742 therein. The side walls 2754 can also include a back cutout 2762 which can align with the air inlet opening 2740 so that air is allowed to flow enter the air pathway 2738 by passing through the back cutout 2762 and through the air inlet opening 2740.

A shaft 2764 can extend downward from the base wall 2752 of the lower housing member 2722, and the shaft 2764 can have a female end 2766 configured to receive the male end of a syringe (not shown). The female end 2766 can include external threads 2768 configured to mate with internal threads of the syringe for securing the syringe thereto. A fluid pathway 2770 can extend from the opening formed in the female end 2766 up through the shaft 2764. The fluid pathway 2770 can include a fork or branch that divides the fluid pathway 2770 so that a fluid inlet opening 2772 and a fluid outlet opening 2774 are both in fluid communication with the fluid pathway 2770. The shaft 2764 can include an enlarged portion 2776 that is wider than the female end 2766 to accommodate the fork or branch in the fluid pathway 2770.

When the top housing member 2720 is attached to the bottom housing member 2722, the fluid outlet opening 2734 of the upper housing member 2720 can align with the fluid inlet opening 2772 of the lower housing member 2722 such that fluid can flow from the vial, through the fluid pathway 2732, out the fluid outlet opening 2734, in the fluid inlet opening 2772, through the fluid pathway 2770, and into the syringe. Also, the fluid inlet opening 2748 of the upper housing member 2720 can align with the fluid outlet opening 2774 of the lower housing member 2722 such that fluid can flow from the syringe, through the fluid pathway 2770, out the fluid outlet opening 2774, in the fluid inlet opening 2748, through the fluid pathway 2746, and to the target connector portion 2706.

A source check valve 2778 can be disposed between the top housing member 2720 and the lower housing member 2722, and can be configured to allow fluid to flow from the fluid outlet opening 2734 to the fluid inlet opening 2772 while preventing fluid from flowing in the reverse direction. The source check valve 2778 can be a duckbill check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction.

A target check valve 2780 can also be disposed between the top housing member 2720 and the lower housing member 2722, and can be configured to allow fluid to flow from the fluid outlet opening 2774 to the fluid inlet opening 2748 while preventing fluid from flowing in the reverse direction. The target check valve 2780 can be a duckbill check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction.

An air check valve 2782 can be disposed between the base 2736 of the upper housing member 2720 and a side wall 2754 of the lower housing member 2722. The check valve 2782 can be positioned between the back cutout 2762 and the air inlet opening 2740 such that air is permitted to flow from the back cutout 2762 to the air inlet opening 2740, but air and fluid are not allowed to flow out of the air inlet opening 2740. The air check valve 2782 can be a duckbill check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction. In some embodiments, a filter (not shown) can be used in conjunction with or in place of the air check valve 2782. The filter can be placed between, or within one of, the back cutout 2762 and the air inlet opening 2740. The filter can be permeable to air so that air is permitted to enter the air passageway 2738. In some embodiments, the filter can be impermeable to the fluid to prevent fluid from exiting the vial via the air pathway 2738. In some embodiments, a bag (not shown) at least partially disposed within the air passageway 2738 can be used to prevent the air that enters the vial from mixing with the fluid. For

example, the piercing member 2724 can include a bag and can be similar to the piercing member 370 discussed above in connection with FIGS. 5A-D.

FIG. 29A is a perspective view of a check valve 2900 which can be used as the source check valve 2778, the target check valve 2780, and/or the air check valve 2782. In some embodiments, the source check valve 2778, the target check valve 2780, and the air check valve 2782 can each have the same shape and size so that they are interchangeable, thereby reducing the cost (e.g., mold creation) that would be required to produce two or three distinct check valve designs. The check valve 2900 can include a base 2902, which can be cylindrical in shape, although other shapes can also be used. A pair of generally opposing bill members 2904a-b can extend upward from the base 2902. The bill members 2904a-b can abut against one another at their ends furthest from the base 2902 forming a slit 2906 therebetween. In the check valve's 2900 relaxed state, the slit 2906 can be closed as shown in FIGS. 29A-B. The base 2902 can include an opening 2908 in fluid communication with a chamber 2910 formed between portions of the bill members 2904a-b.

FIG. 29C is a cross sectional view of the check valve 2900 in the closed configuration. When the slit 2906 is closed and fluid is directed to the check valve 2900 in the direction that the check valve 2900 is configured to block, as shown in FIG. 29C by fluid flow lines, the resulting pressure applied to the outside surfaces of the bill members forces the slit closed. Thus, as greater pressure is applied, the slit 2906 closes more strongly to prevent fluid flow in the undesired direction. Likewise, when fluid is withdrawn from the fluid chamber 2910, the bill members 2904a-b are also drawn together causing the slit 2906 to seal more tightly. FIG. 29D shows the check valve 2900 in the open configuration as fluid is directed through the check valve 2900 in the desired direction, as shown by fluid lines. When fluid is directed through the opening 2908 and into the chamber 2910, the resulting pressure applied to the inside surfaces of the bill members 2904a-b causes the bill members 2904a-b to move away from one another forcing the slit 2906 to open. Likewise, when fluid is drawn away from the outside surfaces of the bill members 2904a-b (with flow in the opposite direction of the flow lines shown in FIG. 29C), the resulting pressure can pull the bill members 2904a-b apart to open the slit 2906. The check valve 2900 can be formed from silicone or any other suitable resilient material.

Returning now to FIGS. 28A-B, the fluid inlet opening 2772 can be wide enough to receive the duckbill portion of the source check valve 2778, and the fluid inlet opening 2748 can be wide enough to receive the duckbill portion of the target check valve 2780. Thus, in some embodiments, the fluid inlet opening 2772 can be wider than the fluid outlet opening 2774, and the fluid inlet opening 2748 can be wider than the fluid outlet opening 2734. The fluid outlet opening 2734 can include a widened end portion that produces a step 2735. The widened portion and the step 2735 can be configured to receive the base of the source check valve 2778. The step 2735 can have a height that is less than the height of the base of the source check valve 2778 so that the base of the check valve 2778 can be compressed between the top housing member 2720 and the lower housing member 2722 when they are attached. Thus, the compressed base of the check valve 2778 can function to seal off the interface between the fluid outlet opening 2734 and the fluid inlet opening 2772 so that fluid can flow through the check valve 2778 without escaping. This can be particularly advantageous when a chemotherapy drug or other hazardous fluid is transported through the connector 2700. The fluid inlet opening 2748 can also have a widened end portion that creates a step 2749 to receive and compress

the base of the target check valve **2780** to seal the interface between the fluid outlet opening **2774** and the fluid inlet opening **2748**. The air inlet opening **2740** can also include a widened end portion that forms a step **2741** and receives the base of the air check valve **2782** to seal the interface between the back cutout **2762** and the air inlet opening **2740**. In some embodiments, all fluid flow paths through the connector are sealed (e.g., hermetically sealed) such that no fluid (e.g., chemotherapy drugs or other hazardous materials) can escape during operation.

FIG. 30A shows the connector **2700**, a vial **3000**, and a syringe **3050** in an unattached configuration. FIG. 30B shows the connector **2700**, the vial **3000**, and the syringe **3050** in an attached configuration. FIG. 30C shows a front view of the connector **2700**. In FIGS. 30A-C, the connector **2700** is illustrated without the target connector portion **2706**. The vial **3000** can include a body **3002**, and a cap **3004**, with a septum **3006** (hidden from view in FIGS. 30A-B) disposed within the cap **3004**. The vial can include a securing ring **3008** formed on the neck of the body **3002**, and/or the cap **3004** can overhang over the edge of the body **3002** forming a securing step **3010**. The vial **3000** can be similar to the vial **314** described herein or any other medical vial or any other suitable container of fluid. It will be understood that various vial shapes and sizes can be used other than the vials shown herein. For example, the vial **3000** can be much larger than the vials (e.g., **314** or **3000**) shown. Also, in some embodiments, other fluid containers can be used in place the vials shown.

As mentioned above, the connector **2700** can include retaining arms **2730a-b** for securing the vial **3000** to the connector **2700**. The manner of securing the vial **3000** to the connector **2700** will be discussed in greater detail with reference to FIGS. 30A-C. The retainer arms **2730a-b** can be general z-shaped, having a lower portion **2784a-b**, a middle portion **2786a-b**, and an upper portion **2788a-b**. The lower portions **2784a-b** can extend outward from the base **2736** of the upper housing member **2720**. As can best be seen in FIG. 30C, the lower portions **2784a-b** can be slightly curved and can angle upward slightly (e.g., at an angle of at least about 10° and/or no more than about 20°, and in some embodiments at an angle of about 15°, from the horizontal plane). The middle portions **2786a-b** can extend inwardly from the ends of the lower portions **2784a-b** and can angle upward at an angle of at least about 30° and/or no more than about 60°, and in some embodiments by an angle of about 45°, from the horizontal plane. The upper portions **2788a-b** can extend outwardly from the ends of the middle portions and can angle upward at an angle of at least about 30° and/or no more than about 60°, and in some embodiments by an angle of about 45°, from the horizontal plane. In some embodiments, the ends of the upward portions **2788a-b** can be curved as best seen in FIG. 30C. Securing projections **2790a-b** can be located at the junctions between the middle portions **2786a-b** and the upper portions **2788a-b**.

The retaining arms **2730a-b** can be formed of a material and thickness such that the retaining arms can resiliently bend outwardly, causing the distance between the securing projections **2790a-b** to increase. To attach the vial **3000** to the connector **2700**, the vial **3000** can be positioned as shown in FIG. 30A, and the vial **3000** can be pushed toward the connector **2700** such that the piercing member **2724** punctures through the septum **3006** of the vial **3000**. As the cap **3004** of the vial **3000** contacts presses against the top/inner surfaces of the upper portions **2788a-b** of the retainer arms **2730a-b**, the retainer arms **2730a-b** can be flexed away from one another until the cap **3004** slips past the securing projections **2790a-b**, at which point the retaining arms **2730a-b** snap back. When

the retaining arms **2730a-b** snap back, the securing projections **2790a-b** can engage the securing step **3010** on the side of the cap **3004** facing the body **3002** of the vial **3000**. In some embodiments, the vial can be advanced until the securing projections **2790a-b** engages with the securing step **3010** on the cap **3004** (as shown in FIG. 30B) or with the securing ring **3008**. In some embodiments, the retaining arms **2730a-b** can include indentations **2792a-b** that can be configured to receive a portion of the vial body **3002** prevent the vial **3000** from shifting once secured to the connector **2700**. If the securing step **3010** on the cap **3004** engages the securing projections **2790a-b**, the securing ring **3008** can engage the indentations **2792a-b** (as shown in FIG. 30B). If the securing ring **3008** engages the securing projections **2790a-b**, the portion of the vial **3000** where the neck widens to the body **3002** can be received by the indentations **2792a-b**.

As shown in FIG. 30B, the piercing member **2724** can extend into the body **3002** of the vial **3000** such that the fluid extraction aperture **2708** is placed into contact with the fluid inside the vial **3000**. In some embodiments, the slit shape of the fluid extraction aperture **2708** can allow the fluid to remain in contact with the fluid extraction aperture **2708** as the fluid is emptied from the vial **3000**. For example, in some embodiments, a portion of the fluid extraction aperture **2708** does not fully pass through the septum so that when the vial **3000** is nearly empty, the little remaining fluid can still be withdrawn through the fluid extraction aperture **2708**. In some embodiments, at least a portion of the septum of the vial can be thicker than the length of the fluid extraction aperture **2708** so that when the piercing member **2724** is inserted through the septum the fluid extraction aperture **2708** is not in simultaneous communication with both the interior and exterior of the vial.

In some embodiments, the connector can include a slit **2894** that extends through a portion of the base **2736** along a midline between the retainer arms **2730a-b**. The slit **2794** can facilitate the flexing of the retainer arms **2730a-b** so that the slit can widen as the arms **2730a-b** are separated from each other. In some embodiments, the piercing member **2724** can connect to the base **2736** of the upper housing member **2720** within an indentation **2796** formed in the upper surface of the base **2736**. The indentation **2796** can also facilitate the flexing of the retainer arms **2730a-b** because the arms **2730** can flex without directly applying pressure to the piercing member **2708**. In some embodiments, the slit **2794** can extend out from the front and back sides of the indentation **2796**.

With further reference to FIGS. 30A-C, the syringe **3050** can be similar to the syringe **318** discussed above, or any other syringe discussed herein. The syringe **3050** can include a body **3052**, a male luer tip **3054**, and a shroud **3056** surrounding the male luer tip **3054**. Internal threads **3058** can be formed on the inside surface of the shroud **3056** to mate with the external threads **2768** formed on the outside surface of the female end **2766**.

It will be understood that the connector **2700** can be used in connection with an automated fluid transfer system (e.g., system **600**). When attached to a fluid transfer station, the connector **2700** can align with sensors for optically detecting the presence of air in the fluid pathway between the vial **3000** and the syringe **3050** as discussed above in connection with FIGS. 17-19D. With further reference now to FIGS. 30B-C, in some embodiments the connector **2700** can be aligned such that the light (e.g., light **676** or **1924**) passes through the fluid pathway **2770** (hidden from view in FIG. 30C) formed in the shaft **2764** within the region **2798** between the enlarged portion **2776** of the shaft **2764** and the location where the upper end of the syringe shroud **3056** ends when the syringe is

47

attached (e.g., as shown in FIG. 30B). In some embodiments, all or a portion of the lower housing member 2722 can be made from a material that is transparent to the light transmitted through the region 2798. In some embodiments, the entire shaft 2764 or the entire portion of the shaft below the enlarged portion 2776 thereof can be transparent. In some embodiments, the shaft 2764 includes a transparent window portion that covers all or a portion of the region 2798, with the remainder of the lower housing member 2722 being made from a material that is opaque to the light.

FIG. 31A shows a cross sectional view of the connector 2700, the vial 3000, and the syringe 3050 as fluid is drawn through the connector 2700 from the vial 3000 to the syringe 3050. As the plunger (not shown) of the syringe 3050 is withdrawn, fluid can be drawn into the body 3052 of the syringe 3050 from the fluid pathway 2770 formed in the shaft 2764. The fluid pathway 2770 can fork or branch so that both the source check valve 2778 and the target check valve 2780 are exposed to the pressure differential caused by the fluid being withdrawn from the fluid pathway 2770. The slit of the target check valve 2780 closes more tightly as fluid is drawn away from it and towards the syringe 3050. The slit of the source check valve 2778 opens as the fluid is drawn toward the syringe. When the source check valve 2778 opens, fluid can be drawn from the source container (e.g., vial 3000) toward the syringe 3050 to compensate for the pressure differential. Fluid can enter the fluid pathway 2732 via the fluid extraction aperture 2708, and flow through the source check valve 2778, into the fluid pathway 2770, and down into the syringe 3050. As fluid is extracted from the vial 3000, air can be drawn into the vial to compensate for the loss of fluid volume. The air can pass through the back cutout 2762, through the air check valve 2782, through the air pathway 2738, and through the air outlet 2712 into the body 3002 of the vial 3000.

FIG. 31B shows a cross sectional view of the connector 2700, the vial 3000, and the syringe 3050 as fluid is driven through the connector 2700 from the syringe 3050 to the target connector portion 2706 which leads to the IV bag assembly (not shown). As the plunger (not shown) of the syringe 3050 is advanced, fluid can be driven from the body 3052 of the syringe 3050 into the fluid pathway 2770 formed in the shaft 2764. The fluid pathway 2770 can fork or branch so that both the source check valve 2778 and the target check valve 2780 are exposed to the pressure differential caused by the fluid being driven into the fluid pathway 2770. The slit of the source check valve 2778 closes more tightly as fluid is pressed against the outside surfaces of its bill members. The slit of the target check valve 2780 opens as the fluid pushed into its chamber and its bill members are pushed away from each other. When the target check valve 2780 opens, fluid can pass through the target check valve 2780, through the fluid pathway 2746, and into the male end 2744 of the target connector portion 2706. Although not shown in FIG. 31B, it will be understood that the fluid can be driven through the target connector portion 2706 and into an IV bag that is attached thereto.

It will be understood that many variations and modifications can be made to the connector 2700. For example, although the illustrated embodiment is shown having an upper housing member 2720 and a lower housing member 2722, it will be understood that the main housing can be made up of a different number of housing members. Some features that are shown as integrated components can be separately formed, and vice versa. For example, in some embodiments, the retaining arms 2730a-b can be separately formed and attachable to the upper housing member 2720. Also, features

48

and elements that are shown as part of the upper housing member 2720 may, in some embodiments, be formed as part of the lower housing member 2722 and vice versa. For example, female end 2742 that is configured to receive the target connector portion 2706 can be formed as part of the lower housing member 2702. Many other variations are also possible.

FIG. 32A is a perspective view of an embodiment of a fluid transfer module in the form of a connector 3200, which can be similar in many regards to the connector 320 or any other connector disclosed herein. FIG. 32B is another perspective view of the connector 3200. The connector 3200 can be used to transfer fluid from a source container (e.g., a vial) to an intermediate measuring container (e.g., a syringe) and then to a target container (e.g., an IV bag). The connector 3200 can include a source connector portion 3202 configured to interface with the source container (e.g., a vial), an intermediate connector portion 3204 configured to interface with the intermediate measuring container (e.g., a syringe), and a target connector portion 3206 configured to interface with the target container (e.g., an IV bag assembly).

The connector 3200 can function to transfer fluid from the source container to the target container similarly to the connector 320 or the connector 2700 or any other connector disclosed herein. Fluid can be extracted from a vial (not shown) through the fluid extraction aperture 3208, and air can enter the vial via the air inlet 3210 and air outlet 3212 to replace the volume of extracted fluid. The fluid extracted from the vial can be drawn through the connector 3200 and into the syringe (not shown) via the opening 3214 formed in the intermediate connector portion 3204. A source check valve (hidden from view in FIGS. 32A-B) can be configured to allow fluid to flow from the fluid extraction aperture 3208 to the opening 3214 in the intermediate connector portion 3204 while preventing fluid from flowing in the reverse direction back into the vial. The fluid can be driven from the syringe into the connector 3200 via the opening 3214, and the fluid can be directed into the target connector portion 3206 and into an IV bag assembly (not shown) attached to the target connector portion 3206. A target check valve (hidden from view in FIGS. 32A-B) can be configured to allow the fluid to flow from the opening 3214 in the intermediate connector portion 3204 to the target connector portion 3206 while preventing fluid from flowing in the reverse direction.

FIG. 33A is an exploded perspective view of the connector 3200. FIG. 33B is another exploded perspective view of the connector 3200. The connector 3200 can include an upper housing member 3220 and a lower housing member 3222. The upper housing member 3220 can include the source connector portion 3202 of the connector 3200, and the lower housing member 3222 can include the intermediate connector portion 3204 of the connector 3200.

The upper housing member 3220 can include a piercing member 3224 made up of an elongate substantially cylindrical shaft 3226 and a pointed tip 3228. The piercing member 3224 can be configured to pierce the septum of a vial (not shown) when the vial is attached thereto. The piercing member 3224 can include a fluid extraction aperture 3208 formed on one side thereof. The fluid extraction aperture can be a slit that extends from near the end of the pointed tip 3228 down onto the shaft 3226, although openings of other shapes can also be used. The piercing member 3224 can also include an air outlet 3212 that allows air to enter the vial as fluid is extracted therefrom to equalize the pressure differential caused by the extraction of fluid. The air outlet 3212 can receive air from an air pathway 3238a that extends through

the shaft 3226 and through the base 3236 and to an air inlet opening 3240 formed in the base 3236 of the upper housing 3220.

The upper housing member 3220 can include a male end 3242 configured to receive a female end 3244 of the target connector portion 3206. The target connector portion 3206 can be similar to the other target connector portions described herein (e.g., 338), the disclosure of which applies also to the target connector portion 3206. In the illustrated embodiment, the target connector portion can include the female end 3244 of the interface while the top housing member can include the male end 3242 thereof. Indeed, any suitable interface for securing the target connector portion 3206 to the upper housing member 3202 can be used. The male end 3242 can be secured to the female end 3244 by applying a plastic welding adhesive (such as Dichloromethane) to the outer surface of the male end 3242 and/or to the inner surface of the female end 3244 before insertion. The Dichloromethane can chemically weld the outer surface of the male end 3242 to the inner surface of the female end 3244. Other methods can be used to connect the male end 3242 to the female end 3244, such as sonic welding, threading, adhesives, etc. In some embodiments, the connection between the male end 3242 and the female end 3244 is hermetically sealed, and in some embodiments includes a sealing member (not shown), such as an O-ring, to provide the hermetic seal. A fluid pathway 3246 can extend from the opening in the male end 3242 to a fluid inlet opening 3248 formed in the bottom surface of the base 3236 of the upper housing member 3220.

The lower housing member 3222 can include a base 3250 configured to mate with the base 3236 of the upper housing member 3220. The base 3236 of the upper housing member 3220 can include a lip 3254 on the bottom surface thereof, forming an indentation. The periphery of the top surface of the base 3250 of the lower housing member 3222 can be configured to contact the bottom surface of the lip 3254 when attached. The upper housing member 3220 can be secured to the lower housing member 3222 using an adhesive, or plastic welding material, or sonic welding, or a snap-fit, or any other suitable technique.

The lower housing member 3222 can include an air inlet 3210 and an air outlet opening 3262 with a fluid pathway 3238b extending therebetween. A shaft 3264 can extend downward from the base 3250 of the lower housing member 3222, and the shaft 3264 can have a female end 3266 configured to receive the male end of a syringe (not shown). The female end 3266 can include external threads 3268 configured to mate with internal threads of the syringe for securing the syringe thereto. A fluid pathway 3270 can extend from the opening formed in the female end 3266 up through the shaft 3264. The fluid pathway 3270 can include a channel 3271 that diverts from the main flow path. Thus the fluid pathway 3270 can provide a fluid inlet opening 3272 and a fluid outlet opening 3274.

When the top housing member 3220 is attached to the bottom housing member 3222, the fluid outlet opening 3234 of the upper housing member 3220 can align with the fluid inlet opening 3272 of the lower housing member 3222 such that fluid can flow from the vial, through the fluid pathway 3232, out the fluid outlet opening 3234, in the fluid inlet opening 3272, through the fluid pathway 3270, and into the syringe. Also, the fluid inlet opening 3248 of the upper housing member 3220 can align with the fluid outlet opening 3274 of the lower housing member 3222 such that fluid can flow from the syringe, through the fluid pathway 3270, out the fluid outlet opening 3274, in the fluid inlet opening 3248, through the fluid pathway 3246, and to the target connector portion

3206. Also, the air outlet opening 3262 can align with the air inlet opening 3240 so that air is allowed to enter through the air inlet 3210, flow through the air pathway 3238b, out the air outlet opening 3262, in the air inlet opening 3240, through the air pathway 3238a, through the air outlet 3212 and into the vial.

A check valve assembly 3277 can be disposed between the top housing member 3220 and the lower housing member 3222. The check valve assembly 3277 can include a base which can be shaped to fit into the indentation formed by the lip 3254. The check valve assembly 3277 can include a source check valve 3278 configured to allow fluid to flow from the fluid outlet opening 3234 to the fluid inlet opening 3272 while preventing fluid from flowing in the reverse direction. The source check valve 3278 can be a dome valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction.

The check valve assembly 3277 can include a target check valve 3280 configured to allow fluid to flow from the fluid outlet opening 3274 to the fluid inlet opening 3248 while preventing fluid from flowing in the reverse direction. The target check valve 3280 can be a domed check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction.

The check valve assembly 3277 can include an air check valve 3282 configured such that air is permitted to flow from the air outlet 3262 to the air inlet opening 3240, but air and fluid are not allowed to flow out of the air inlet opening 3240. The air check valve 3282 can be a domed check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction. In some embodiments, a filter (not shown) can be used in conjunction with or in place of the air check valve 3282. The filter can be placed in or near the air inlet, or within the air pathways 3238a-b. The filter can be permeable to air so that air is permitted to enter the air passageway 3238a-b. In some embodiments, the filter can be impermeable to the fluid to prevent fluid from exiting the vial via the air pathway 3238a-b. In some embodiments, a bag (not shown) at least partially disposed within the air passageway 3238a can be used to prevent the air that enters the vial from mixing with the fluid. For example, the piercing member 3224 can include a bag and can be similar to the piercing member 370 discussed above in connection with FIGS. 5A-D.

Although the domed check valves 3278, 3280, 3282 are shown as being interconnected by the base 3279, it will be understood that the domed check valves 3278, 3280, 3282 can be separately formed. A domed check valve can include a dome having a convex side and a concave side. One or more slits 3281 can be formed in the dome. Although a single slit is shown in the illustrated embodiment, it will be understood that two crossing slits, or various other slit configurations can be used. In the domed check valve's relaxed state, the slit can be closed.

When the slit 3281 is closed and fluid is directed to the check valve 3278, 3280, 3282 in the direction that the check valve 3278, 3280, 3282 is configured to block, the resulting pressure that pushes on the convex side forces the slit 3281 closed. Thus, as greater pressure is applied, the slit 3281 closes more strongly to prevent fluid flow in the undesired direction. Likewise, when fluid is withdrawn from the concave side, the slit 3281 is sealed more tightly. When fluid is pushed toward the concave side, the resulting pressure causes the dome to flex outwardly such that the slit 3281 opens.

Likewise, when fluid is drawn away from the convex side, the resulting pressure can pull the dome members such that they flex outwardly and the slit 3281 opens. The check valve assembly 3277 can be formed from silicone or any other suitable resilient material.

With further reference to FIGS. 33A-B, the fluid inlet opening 3272 can be wide enough to receive the dome portion of the source check valve 3278, and the fluid inlet opening 3248 can be wide enough to receive the dome portion of the target check valve 3280. Thus, in some embodiments, the fluid inlet opening 3272 can be wider than the channel 3271 that functions as the fluid outlet opening 3274, and the fluid inlet opening 3248 can be wider than the fluid outlet opening 3234. The indentation formed by the lip 3254 can have a height that is less than the height of the base 3279 of the check valve assembly 3277 so that the base 3279 can be compressed between the top housing member 3220 and the lower housing member 3222 when they are attached. Thus, the compressed base 3279 of the check valve assembly 3277 can function to seal off the interfaces between the upper housing member 3220 and the lower housing member 3222 so that fluid can flow therethrough without escaping. This can be particularly advantageous when a chemotherapy drug or other hazardous fluid is transported through the connector 3200. In some embodiments, all fluid flow paths through the connector 3200 are sealed (e.g., hermetically sealed) such that no fluid (e.g., chemotherapy drugs or other hazardous materials) can escape during operation.

FIG. 34A shows a cross sectional view of the connector 3200, the vial 3000, and the syringe 3050 as fluid is drawn through the connector 3200 from the vial 3000 to the syringe 3050. As the plunger (not shown) of the syringe 3050 is withdrawn, fluid can be drawn into the body 3052 of the syringe 3050 from the fluid pathway 3270 formed in the shaft 3264. The fluid can be drawn in from the pathway 3270 including the channel 3271 so that both the source check valve 3278 and the target check valve 3280 are exposed to the pressure differential caused by the fluid being withdrawn from the fluid pathway 3270. The slit of the target check valve 3280 closes more tightly as fluid is drawn away from it and towards the syringe 3050. The slit of the source check valve 3278 opens as the fluid is drawn toward the syringe. When the source check valve 3278 opens, fluid can be drawn from the source container (e.g., vial 3000) toward the syringe 3050 to compensate for the pressure differential. Fluid can enter the fluid pathway 3232 via the fluid extraction aperture 3208, and flow through the source check valve 3278, into the fluid pathway 3270, and down into the syringe 3050. As fluid is extracted from the vial 3000, air can be drawn into the vial 3000 to compensate for the loss of fluid volume. The air can pass through the air inlet 3210, through the air pathway 3238b, through the air check valve 3282, through the air pathway 3238a, and through the air outlet 3212 into the body 3002 of the vial 3000.

FIG. 34B shows a cross sectional view of the connector 3200, the vial 3000, and the syringe 3050 as fluid is driven through the connector 3200 from the syringe 3050 to the target connector portion 3206 which leads to the IV bag assembly (not shown). As the plunger (not shown) of the syringe 3050 is advanced, fluid can be driven from the body 3052 of the syringe 3050 into the fluid pathway 3270 formed in the shaft 3264. The fluid can enter the channel 3271 so that both the source check valve 3278 and the target check valve 3280 are exposed to the pressure differential caused by the fluid being driven into the fluid pathway 3270. The slit of the source check valve 3278 closes more tightly as fluid is pressed against the convex surface of its dome. The slit of the

target check valve 3280 opens as the fluid pushed against the concave surface of its dome. When the target check valve 3280 opens, fluid can pass through the target check valve 3280, through the fluid pathway 3246, and into the female end 3244 of the target connector portion 3206. Although not shown in FIG. 34B, it will be understood that the fluid can be driven through the target connector portion 3206 and into an IV bag that is attached thereto.

It will be understood that the connector 3200 can be used in connection with an automated fluid transfer system (e.g., system 600). When attached to a fluid transfer station, the connector 3200 can align with sensors for optically detecting the presence of air in the fluid pathway between the vial 3000 and the syringe 3050 as discussed above in connection with FIGS. 17-19D. With further reference now to FIGS. 34A-B, in some embodiments the connector 3200 can be aligned such that the light (e.g., light 676 or 1924) passes through the fluid pathway 3270 formed in the shaft 3264 within the region 3298 above the location where the upper end of the syringe shroud 3056 ends when the syringe 3050 is attached. In some embodiments, all or a portion of the lower housing member 3222 can be made from a material that is transparent to the light transmitted through the region 3298. In some embodiments, the entire shaft 3264 can be transparent. In some embodiments, the shaft 3264 includes a transparent window portion that covers all or a portion of the region 3298, with the remainder of the lower housing member 3222 being made from a material that is opaque to the light.

It will be understood that many variations and modifications can be made to the connector 3200. For example, although the illustrated embodiment is shown having an upper housing member 3220 and a lower housing member 3222, it will be understood that the main housing can be made up of a different number of housing members. Also, features and elements that are shown as part of the upper housing member 3220 may, in some embodiments, be formed as part of the lower housing member 3222 and vice versa.

FIG. 35A is a perspective view of an embodiment of a connector 3500, which can be similar in many regards to the connector 350 or any other connector disclosed herein. FIG. 35B is another perspective view of the connector 3500. The connector 3500 can be used to transfer fluid from a source container (e.g., a vial) to an intermediate measuring container (e.g., a syringe) and then to a target container (e.g., an IV bag). The connector 3500 can include a source connector portion 3502 configured to interface with the source container (e.g., a vial), an intermediate connector portion 3504 configured to interface with the intermediate measuring container (e.g., a syringe), and a target connector portion 3506 configured to interface with the target container (e.g., an IV bag assembly).

The connector 3500 can function to transfer fluid from the source container to the target container similarly to the connector 350 or the connector 2700 or any other connector disclosed herein. Fluid can be extracted from a vial (not shown) through the fluid extraction aperture 3508, and air can enter the vial via the air inlet 3510 and air outlet 3512 to replace the volume of extracted fluid. The fluid extracted from the vial can be drawn through the connector 3500 and into the syringe (not shown) via the opening 3514 formed in the intermediate connector portion 3504. A source check valve (hidden from view in FIGS. 35A-B) can be configured to allow fluid to flow from the fluid extraction aperture 3508 to the opening 3514 in the intermediate connector portion 3504 while preventing fluid from flowing in the reverse direction back into the vial. The fluid can be driven from the syringe into the connector 3500 via the opening 3514, and the fluid can be directed into the target connector portion 3506 and into

an IV bag assembly (not shown) attached to the target connector portion **3506**. A target check valve (hidden from view in FIGS. **35A-B**) can be configured to allow the fluid to flow from the opening **3514** in the intermediate connector portion **3504** to the target connector portion **3506** while preventing fluid from flowing in the reverse direction.

FIG. **36A** is an exploded perspective view of the connector **3500**. FIG. **36B** is another exploded perspective view of the connector **3500**. The connector **3500** can include an upper housing member **3520** and a lower housing member **3522**. The upper housing member **3520** can include the source connector portion **3502** of the connector **3500**, and the lower housing member **3522** can include the intermediate connector portion **3504** of the connector **3500**.

The upper housing member **3520** can include a piercing member **3524** made up of an elongate substantially cylindrical shaft **3526** and a pointed tip **3528**. The piercing member **3524** can be configured to pierce the septum of a vial (not shown) when the vial is attached thereto. The upper housing member **3220** can include retaining arms **3230a-b** configured to secure the vial to the connector **2700** in a manner similar to that described in connection with the retaining arms **2730a-b**. The piercing member **3524** can include a fluid extraction aperture **3508** formed on one side thereof. The fluid extraction aperture can be a slit that extends from near the end of the pointed tip **3528** down onto the shaft **3526**, although openings of other shapes can also be used. The piercing member **3524** can also include an air outlet **3512** that allows air to enter the vial as fluid is extracted therefrom to equalize the pressure differential caused by the extraction of fluid. The air outlet **3512** can receive air from an air pathway **3538a** that extends through the shaft **3526** and through the base **3536** and to an air inlet opening **3540** formed in the base **3536** of the upper housing **3520**.

The upper housing member **3520** can include a female end **3542** configured to receive a male end **3544** of the target connector portion **3506**. The target connector portion **3506** can be similar to the other target connector portions described herein (e.g., **338**), the disclosure of which applies also to the target connector portion **3506**. Any suitable interface for securing the target connector portion **3506** to the upper housing member **3502** can be used. The female end **3542** can be secured to the male end **3544** by applying a plastic welding adhesive (such as Dichloromethane) to the outer surface of the male end **3544** and/or to the inner surface of the female end **3542** before insertion. The Dichloromethane can chemically weld the outer surface of the male end **3544** to the inner surface of the female end **3542**. Other methods can be used to connect the male end **3544** to the female end **3542**, such as sonic welding, threading, adhesives, etc. In some embodiments, the connection between the male end **3544** and the female end **3542** is hermetically sealed, and in some embodiments includes a sealing member (not shown), such as an O-ring, to provide the hermetic seal. A fluid pathway **3546** can extend from the opening in the female end **3542** to a fluid inlet opening **3548** formed in the bottom surface of the base **3536** of the upper housing member **3520**.

The lower housing member **3522** can include a chamber **3550** enclosed by a base wall **3252** and by side walls **3254** and can have an open top. The chamber **3250** can be configured to receive the base **3536** of the upper housing member **2720** when the top housing member **3520** is secured to the bottom housing member **3522**. The side walls **3554** can include a lip **3556** near the top thereof which can be configured to mate with corresponding slots **3558** formed in the upper portion of the base **3536** for provide a snap-fit connection between the top housing member **3520** and the bottom housing member

3522. It will be understood that the top housing member **3520** can be secured to the bottom housing member **3522** using various other techniques including an adhesive, sonic welding, a friction-fit, or any other suitable manner. The side walls **3554** of the lower housing member **3522** can include a front cutout **3560** configured to receive a portion of the female end **3542** therein.

The lower housing member **3522** can include an air inlet **3510** and an air outlet opening **3562** with a fluid pathway **3538b** extending therebetween. A shaft **3564** can extend downward from the base wall **3552** of the lower housing member **3522**, and the shaft **3564** can have a female end **3566** configured to receive the male end of a syringe (not shown). The female end **3566** can include external threads **3568** configured to mate with internal threads of the syringe for securing the syringe thereto. A fluid pathway **3570** can extend from the opening formed in the female end **3566** up through the shaft **3564**. The fluid pathway **3570** can include a fork or branch that divides the fluid pathway **3570** so that a fluid inlet opening **3572** and a fluid outlet opening **3574** are both in fluid communication with the fluid pathway **3570**.

When the top housing member **3520** is attached to the bottom housing member **3522**, the fluid outlet opening **3534** of the upper housing member **3520** can align with the fluid inlet opening **3572** of the lower housing member **3522** such that fluid can flow from the vial, through the fluid pathway **3532**, out the fluid outlet opening **3534**, in the fluid inlet opening **3572**, through the fluid pathway **3570**, and into the syringe. Also, the fluid inlet opening **3548** of the upper housing member **3520** can align with the fluid outlet opening **3574** of the lower housing member **3522** such that fluid can flow from the syringe, through the fluid pathway **3570**, out the fluid outlet opening **3574**, in the fluid inlet opening **3548**, through the fluid pathway **3546**, and to the target connector portion **3506**. Also, the air outlet opening **3562** can align with the air inlet opening **3540** so that air is allowed to enter through the air inlet **3510**, flow through the air pathway **3538b**, out the air outlet opening **3562**, in the air inlet opening **3540**, through the air pathway **3538a**, through the air outlet **3512** and into the vial.

A check valve assembly **3577** can be disposed between the top housing member **3520** and the lower housing member **3522**. The check valve assembly **3577** can include a source check valve **3578** configured to allow fluid to flow from the fluid outlet opening **3534** to the fluid inlet opening **3572** while preventing fluid from flowing in the reverse direction. The source check valve **3578** can be a flap check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction.

The check valve assembly **3577** can include a target check valve **3580** configured to allow fluid to flow from the fluid outlet opening **3574** to the fluid inlet opening **3548** while preventing fluid from flowing in the reverse direction. The target check valve **3580** can be a flap check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction.

The check valve assembly **3577** can include an air check valve **3582** configured such that air is permitted to flow from the air outlet **3562** to the air inlet opening **3540**, but air and fluid are not allowed to flow out of the air inlet opening **3540**. The air check valve **3582** can be a flap check valve as shown in the illustrated embodiment, or any other form of check valve capable of allowing fluid to flow in one direction while preventing fluid flow in the opposite direction. In some embodiments, a filter (not shown) can be used in conjunction

55

with or in place of the air check valve **3582**. The filter can be placed in or near the air inlet **3510**, or within the air pathway **3538a-b**. The filter can be permeable to air so that air is permitted to enter the air pathway **3538a-b**. In some embodiments, the filter can be impermeable to the fluid to prevent fluid from exiting the vial via the air pathway **3538a-b**. In some embodiments, a bag (not shown) at least partially disposed within the air pathway **3538a** can be used to prevent the air that enters the vial from mixing with the fluid. For example, the piercing member **3524** can include a bag and can be similar to the piercing member **370** discussed above in connection with FIGS. 5A-D.

FIG. 37 is a perspective view of a check valve assembly **3700** which can be used as the check valve assembly **3577** discussed herein. The check valve assembly **3577** can include a base **3702** with a right opening **3704**, a central opening **3706**, and a left opening **3708** formed therethrough. A series of raised ridges **3722a** can outline the openings **3704**, **3706**, **3708** on the top side of the base **3702**, and a series of raised ridges **3722b** can outline the openings **3704**, **3706**, **3708** on the bottom side of the base **3702**. A right divider **3710** can divide the right opening **3704** from the central opening **3706**. A left divider **3712** can divide the left opening **3708** from the central opening **3706**.

A right flap **3714** can extend from the right divider **3710** into the right opening **3704**. The right flap **3714** can be sized so as to cover a substantial portion of the right opening **3704** but leaving a narrow open area surrounding the right flap **3714**. A left flap **3716** can extend from the left divider **3712** into the left opening **3708**. The left flap **3716** can be sized so as to cover a substantial portion of the left opening **3708** but leaving a narrow open area surrounding the left flap **3716**. A first central flap **3718** can extend from the right divider **3710** into the central opening **3706**. A second central flap **3720** can extend from the left divider **3712** into the central opening **3706**. The first and second central flaps **3718**, **3720** can be configured to fill a substantial portion of the central opening **3706** but leaving a narrow open area surrounding the first and second central flaps **3718**, **3720**.

The flaps **3714**, **3716**, **3718**, **3720** can resiliently deform to open a fluid pathway. The flaps **3714**, **3716**, **3718**, **3720** are shown in FIG. 37 in relaxed positions. However, if a force (e.g., fluid pressure) is applied to one side of a flap **3714**, **3716**, **3718**, **3720**, the flap **3714**, **3716**, **3718**, **3720** can be displaced in the direction of the applied force. In some embodiments, the flaps **3714**, **3716**, **3718**, **3720** can pivot or hinge on the dividers **3710**, **3712** and/or the flaps **3714**, **3716**, **3718**, **3720** themselves can bend to assume a curved shape. The manner in which the flaps **3714**, **3716**, **3718**, **3720** operation as check valves will be described in greater detail below.

In some embodiments, the check valve assembly **3700** can be symmetrical across the x-y plane, the x-z plane, and/or the y-z plane. This symmetry can facilitate assembly of the connector because the check valve assembly **3700** cannot be inserted backwards or upside-down.

Returning now to FIGS. 36A-B, the check valve assembly **3577** can include a source check valve **3578** (e.g., second central flap **3720**), and a target check valve **3580** (e.g., right flap **3714**), and an air check valve **3582** (e.g., left flap **3716**). In some embodiments, the check valve assembly **3577** can include an extra flap **3583** (e.g., first central flap **3718**) that does not function as a check valve. The extra flap **3581** can be included to maintain the symmetry of the check valve assembly **3577** to simplify assembly of the connector **2500**.

With further reference to FIGS. 33A-B, the fluid inlet opening **3572** can be wide enough to allow the source check valve **3578** to swing open, but the fluid outlet opening **3534**

56

can fit flush against the flap of the source check valve **3578**, thereby allowing the flap of the source check valve **3578** to open only in the direction toward the fluid pathway **2770**. The fluid inlet opening **3548** can be wide enough to allow the target check **3580** valve to swing open, but the fluid outlet opening **3574** can fit flush against the flap of the target check valve **3580**, thereby allowing the flap of the target check valve **3580** to open only in the direction toward the fluid pathway **3546**. The air inlet opening **3540** can be wide enough to allow the air check valve **3582** to swing open, but the air outlet opening **3562** can fit flush against the flap of the air check valve **3582**, thereby allowing the flap of the air check valve **3582** to open only in the direction toward the fluid pathway **3538a**. The functionality of the check valves **3578**, **3580**, and **3582** can also be seen in FIGS. 38A-B which will be discussed below.

The height of the base **3702** and/or ridges **3722a-b** of the check valve assembly **2577** can be configured such that the base **3702** and/or ridges **3722a-b** are compressed between the top housing member **3520** and the lower housing member **3522** when they are attached. Thus, the compressed base **3702** and/or ridges **3722a-b** of the check valve assembly **2577** can function to seal off the interfaces between the upper housing member **3520** and the lower housing member **3522** so that fluid can flow therethrough without escaping. This can be particularly advantageous when a chemotherapy drug or other hazardous fluid is transported through the connector **3500**. In some embodiments, all fluid flow paths through the connector **3500** are sealed (e.g., hermetically sealed) such that no fluid (e.g., chemotherapy drugs or other hazardous materials) can escape during operation.

FIG. 38A shows a cross sectional view of the connector **3500**, the vial **3000**, and the syringe **3050** as fluid is drawn through the connector **3500** from the vial **3000** to the syringe **3050**. As the plunger (not shown) of the syringe **3050** is withdrawn, fluid can be drawn into the body **3052** of the syringe **3050** from the fluid pathway **3570** formed in the shaft **3564**. Because the fluid pathway **3570** forks or branches, both the source check valve **3578** and the target check valve **3580** are exposed to the pressure differential caused by the fluid being withdrawn from the fluid pathway **3570**. The pressure differential caused by the fluid being withdrawn from the fluid pathway **3570** pulls the flap of the target check valve **3580** more firmly closed against the base wall **3552** because the fluid outlet opening **3574** is not wide enough to accommodate the flap. The pressure differential can pull the flap of the source check valve **3578** open. When the source check valve **3578** opens, fluid can be drawn from the source container (e.g., vial **3000**) toward the syringe **3050** to compensate for the pressure differential. Fluid can enter the fluid pathway **3532** via the fluid extraction aperture **3508**, and flow past the source check valve **3578**, into the fluid pathway **3570**, and down into the syringe **3050**. The extra flap **3583** can also be pulled down into the fluid inlet opening **3572** toward the fluid pathway **3570**. In some embodiments, the extra flap **3583** does not function as a check valve and does not substantially affect the flow of fluid in either the relaxed or deformed configuration. In some embodiments, the extra flap **3583** can be omitted. As fluid is extracted from the vial **3000**, air can be drawn into the vial **3000** to compensate for the loss of fluid volume. The air can pass through the air inlet **3510**, through the air pathway **3538b**, past the air check valve **3582**, through the air pathway **3538a**, and through the air outlet **3512** into the body **3002** of the vial **3000**.

FIG. 38B shows a cross sectional view of the connector **3500**, the vial **3000**, and the syringe **3050** as fluid is driven through the connector **3500** from the syringe **3050** to the

57

target connector portion **3506** which leads to the IV bag assembly (not shown). As the plunger (not shown) of the syringe **3050** is advanced, fluid can be driven from the body **3052** of the syringe **3050** into the fluid pathway **3570** formed in the shaft **3564**. The fluid pathway **3570** can fork or branch so that both the source check valve **3578** and the target check valve **3580** are exposed to the pressure differential caused by the fluid being driven into the fluid pathway **3570**. The pressure differential caused by the fluid being driven into the fluid pathway **3570** can push the flap of the source check valve **3578** more firmly closed against the bottom surface of the base **2536** because the fluid outlet opening **3534** is not wide enough to accommodate the flap. The flap of the target check valve **3580** can swing open as the fluid pushed against the flap. When the target check valve **3580** opens, fluid can flow past the target check valve **3580**, through the fluid pathway **3546**, and into the male end **3544** of the target connector portion **3506**. Although not shown in FIG. **38B**, it will be understood that the fluid can be driven through the target connector portion **3506** and into an IV bag that is attached thereto.

It will be understood that the connector **3500** can be used in connection with an automated fluid transfer system (e.g., system **600**). When attached to a fluid transfer station, the connector **3500** can align with sensors for optically detecting the presence of air in the fluid pathway between the vial **3000** and the syringe **3050** as discussed above in connection with FIGS. **17-19D**. With further reference now to FIGS. **38A-B**, in some embodiments the connector **3500** can be aligned such that the light (e.g., light **676** or **1924**) passes through the fluid pathway **3570** formed in the shaft **3564** within the region **3598** above the location where the upper end of the syringe shroud **3056** ends when the syringe **3050** is attached. In some embodiments, all or a portion of the lower housing member **3522** can be made from a material that is transparent to the light transmitted through the region **3598**. In some embodiments, the entire shaft **3564** can be transparent. In some embodiments, the shaft **3564** includes a transparent window portion that covers all or a portion of the region **3598**, with the remainder of the lower housing member **3522** being made from a material that is opaque to the light.

It will be understood that many variations and modifications can be made to the connector **3500**. For example, although the illustrated embodiment is shown having an upper housing member **3520** and a lower housing member **3522**, it will be understood that the main housing can be made up of a different number of housing members. Also, features and elements that are shown as part of the upper housing member **3520** may, in some embodiments, be formed as part of the lower housing member **3522** and vice versa.

Several connectors for transferring fluid are described herein (e.g., connectors **320**, **2600**, **2700**, **3200**, **3500**, **3910**). It will be understood that many of the features described in connection with one connector can also be applied to the other connectors disclosed herein. Many components of the connectors can be interchangeable with corresponding components of the other connectors. For example, The connectors **2700** and **3500** are shown as having retaining arms for securing a vial thereto, and the retaining arms can similarly be incorporated into the other connectors (e.g., **320** or **3200**). Indeed, in some embodiments, the retaining arms can be removably attachable and can slide over the piercing member and snap into place into a groove formed in the base of the shaft of the piercing member (see FIG. **32A**). Each of the connectors can be modified to incorporate the check valve types disclosed in connection with each of the other connectors. In some embodiments, a single connector can use dif-

58

ferent check valve types for different check valves. One possible configuration is to use a series of three duckbill check valves (e.g., as shown in connector **2700**) but integrated into a single check valve assembly and oriented similar to the check valve assembly of the connector **3200**. Many other modifications are possible.

FIG. **39** is a perspective view of another example embodiment of a fluid transfer system **3900**. The fluid transfer station **3900** can be similar to, or the same as, fluid transfer systems **100** or **600** or any other fluid transfer system discussed herein. Thus, the discussion associated with many features of other fluid transfer systems described herein is also applicable to the fluid transfer system **3900**, even when not specifically identified.

The fluid transfer system can include a main housing **3902** that supports two transfer stations **3904a-b**, although any other suitable number of transfer stations can be used (e.g. one, three, four, five, or more transfer stations). The transfer stations **3904a-b** can be similar to, or the same as, the transfer stations **604a-f** discussed above. Although only transfer station **604a** is discussed in further detail below, it should be understood that the transfer station **604b** can be the same as transfer station **604a**, or the transfer stations **604a-b** can vary (e.g., having different sized syringes).

The transfer station **3904a** can be configured to receive a fluidics assembly **3906** in a manner similar to that described in connection with transfer station **604a**. The fluidics assembly **3906** can include a vial (not shown in FIG. **39**), a vial adapter **3908**, a fluid transfer module or connector **3910**, a syringe **3912**, and an IV bag assembly **3914** (partially shown in FIG. **39**). The transfer station can be configured to secure the syringe **3912** and/or connector **3910** using, for example, a top connector **3916**, a middle connector **3918**, and an end piece **3920**. The transfer station **3904a** can include a motor (inside the housing **3902**) to cause the end piece **3920** to move with respect to the middle connector **3918**, thus withdrawing or advancing the plunger of the syringe **3912**. In some embodiments, the motor can be a high precision stepping motor able to withdraw the plunger of the syringe **3912** by a precise distance, thereby facilitating precision fluid transfer. In some embodiments, the system **3900** can transfer amounts of fluid in increments within the range of approximately 0.05 milliliters to approximately 0.3 milliliters. In some embodiments, the system **3900** can transfer amounts of fluid in increments of about 0.1 milliliters. In some embodiments, the system **3900** can transfer fluid at a rate in the range of about 10 to 70 milliliters per minute for each transfer station. In some embodiments, the rate can be about 30 milliliters per minute for each fluid transfer station. In some embodiments, the system **3900** can transfer fluid with an error rate in the range of about 0% to about 8% when transferring a volume of more than 1 milliliter. In some embodiments, the error rate can be about 3%.

In some embodiments fluid transfer station **3904a** can include a compatibility mechanism configured to ensure that an approved connector is used, to provide reliable accurate fluid transfer. The compatibility mechanism can be a mounting feature (e.g., of the top connector **3916**) that is configured specifically to fit with a portion of the connector **3910**. In some embodiments, the fluid transfer module or connector **3910** can be a single-use, disposable portion. The fluid transfer module **3910** can be provided with instructions to the user for inserting the fluid transfer module **3910** into the electronically controlled fluid dispensing system to properly position and align the various components to allow for fluid transfer and safety features. The fluid transfer module **3910** also can be provided with instructions to the user for disconnecting the

fluid transfer module **3910** after fluid transfer is completed. In some embodiments, the user instructions can include information indicating that the fluid transfer module should be disposed of in a biohazard receptacle after a single use.

The fluid transfer station **3904a** can include a tray **3922** to support the IV bag assembly **3914**. The tray **3922** can be similar to, or the same as the tray **2272** described above. In some embodiments, the tray **3922** can be secured to the top connector **3916** or other portion of the housing **3902** using screws or the tray **3922** can be inserted into a slot. Other supports can be used. In some embodiments, the tray **3922** can pivot down when not in use, as will be discussed in greater detail below.

An electronically controlled fluid dispensing system, such as the fluid transfer system **3900** can include a power switch **3926**, and various input and/or output ports **3928** for connecting external devices (e.g., a keypad, touchscreen, controller, printer, barcode scanner, monitor, or computer). In some embodiments a foot pedal can connect to one of the ports **3928**. The foot pedal can include a button or switch to start and stop the fluid transfer process. The housing **3902** can have support feet **3930** extending therefrom, and handles **3932**.

FIG. **40** is a perspective view of the fluidics assembly **3906** in an assembled configuration. FIG. **41** is a perspective exploded view of the fluidics assembly **3906** from a different angle than that shown in FIG. **40**. The fluid assembly **3906** can be used to transfer precise amounts of fluid from the vial **3907** to the IV bag **3914**. The fluidics assembly **3906** includes a vial **3907**, a vial adapter **3908** configured to provide fluid communication with the fluid (e.g., chemotherapy drug or other medication) contained within the vial, a syringe **3912**, an IV bag assembly **3914**, and a connector **3910** for directing fluid from the vial adapter **3908** into the syringe **3912** and from the syringe toward the IV bag assembly. In some embodiments, the fluidics assembly **3906** can have features similar to, or the same as, those of the other fluidics systems disclosed. In some embodiments, the fluidics assembly **3906** can be configured to allow the vial **3907** and vial adapter **3908** to be replaced (e.g., when the vial runs out of fluid) without replacing the connector **3910** or syringe **3912**. Unlike many of the connectors disclosed herein, in the fluidics assembly **3906**, air enters the vial **3907** via the vial adapter **3908** rather than through the connector **3910**.

FIG. **42** is a perspective view showing the vial adapter **3908** and the vial **3907** in a separated configuration, such as before the vial **3907** is attached to the vial adapter **3908**. The vial adapter can have a top portion **3940** that is similar to, or the same as, the top of the connector **2700**, the connector **3500**, or any of the other connectors described as being able to access fluid in a vial (or bag or other fluid source container). For example, the top portion **3940** can include a spike **3942** configured to pierce the septum on the cap of the vial **3907** and arms **3942** to retain the vial **3907** onto the vial adapter **3908**.

Opposite the upper portion **3940**, the vial adapter can include a connector, which can be, for example, a female connector **3944**. The connector **3944** can be, for example, a version of the Clave® connector manufactured by ICU Medical, Inc., of San Clemente, Calif. Various embodiments of a connector of this type are described in the '866 Patent. The female connector **3944** can seal the end of the vial adapter **3908** such that no fluid is allowed to escape from the vial adapter **3908** until a male connector is attached to the female connector **3944**. It should be understood that in many embodiments discussed herein, the male and female connectors can be switched. For example, the vial adapter **3908** can include a male connector which is configured to mate with a female connector on the connector **3910**.

The vial adapter **3908** can include an air intake channel **3946** configured to direct air into the vial **3907** to compensate for fluid removed from the vial **3907** to reduce the pressure differential. The air intake channel **3946** can include a filter **3948** configured to allow air to pass through the filter **3948** and toward the vial **3907** while also preventing fluid from passing through the filter. For example, the filter **3948** can include an air permeable but fluid impermeable membrane. The filter **3948** can be a hydrophobic filter. In some embodiments, the vial adapter **3908** can include a check valve in place of or in addition to the filter **3948**. The vial adapter **3908** can also have a bag that is configured to increase in volume while preventing the input air to contact the fluid inside the vial **3907**, similar to the bag **394** discussed above. Thus, the vial **3907** can be vented by a mechanism independent of the connector **3910**.

FIG. **43** is a cross sectional view of the vial **3907** and vial adapter **3908** in an assembled configuration. As shown by the flow lines in FIG. **43**, Air can pass through the filter **3948**, through the air inlet channel **3946**, and into the vial **3907** to compensate for the fluid that is drawn out of the vial **3907** through a fluid channel **3950**. The fluid channel **3950** can pass through the spike **3942**, and down through the female connector **3944** as shown. Although the female connector **3944** is shown in a closed configuration in FIG. **43**, it will be understood that the female connector **3944** can be opened by the first male connector **3964** of the connector **3910** to allow fluid to pass from the vial adapter **3908** to the connector **3910**.

FIG. **44** is a perspective view of the connector **3910**. FIG. **45** is a perspective view of the connector taken from a different angle than the view of FIG. **44**. FIG. **46** is a right-side view of the connector **3910**. FIG. **47** is a back view of the connector **3910**. FIG. **48** is a view of the connector **3910**. FIG. **49** is a top-down view of the connector **3910**. FIG. **50** is a bottom-up view of the connector **3910**. FIG. **51** is a left-side view of the connector **3910**.

The connector **3910** can have features similar to, or the same as, those of the connector **2700** or any other connector disclosed here. The connector **3910** can include an upper housing portion **3960** and a lower housing portion **3962**. A first male connector **3964** can be attached to a female end **3966** of the upper housing portion. A second male connector **3964** can be attached to a female end **3968** of the lower housing portions **3962**. The male connectors **3964**, **3968** can be a version of the Spiros® closeable male connector manufactured by ICU Medical, Inc., of San Clemente, Calif. Various embodiments of connectors of this type are described in the '920 Publication. A syringe interface **3972** can extend down from the bottom of the lower housing portion **3962** to receive the syringe **3912**. A sensor region **3974** can also be positioned at the base of the lower housing portion **3962** and can be configured to allow light to pass through the fluid pathway in the connector **3910** to detect the presence of bubbles, which can indicate that the vial **3907** has run out of fluid. In some embodiments, the surface of the sensor region can be flat to allow light to pass through the wall of the sensor region **3974** at an angle that is perpendicular to the surface, thereby allowing the light to more reliably strike the corresponding sensor.

FIG. **52** is an exploded perspective view of the connector **3910**. FIG. **53** is an exploded perspective view of the connector **3910** taken from a different view than FIG. **52**. The connector **3910** can be similar to the connector **2700** in many respects. However, instead of including a vial adapter built into the upper housing portion, as is the case for the connector **2700**, the connector **3910** includes the first male connector **3964** which is configured to removably interface with the

61

female connector **3944** of the separate vial adapter **3908**. Thus, when the vial **3907** runs out of fluid, the vial **3907** and vial adapter **3908** can be replaced without replacing the connector **3910**, syringe **3912**, or any other part of the fluidics assembly **3906**. This can provide the benefit of reducing the amount of disposable pieces and fluid sent to waste during a vial replacement. Because the vial adapter is not part of the connector **3910**, the connector **3910** also differs from the connector **2700** in that the connector **3910** does not include an air inlet channel or an air check valve. Other connectors which are described herein as having an integrated vial adapter (e.g., the connectors **320**, **3200**, **3500**) can be similarly modified to be compatible with a separate vial adapter.

When the vial **3907**, vial adapter **3908**, connector **3910**, syringe **3912**, and IV bag assembly **3914** are connected, a source fluid pathway can be formed between the vial **3907** and the syringe **3912**, and a target fluid pathway can be formed between the syringe **3912** and the IV bag. The connector **3910** can include a source check valve **3976** positioned in the source fluid pathway to allow fluid to flow from the vial **3907** into the syringe and prevent fluid from flowing back into the vial **3907**. The connector **3910** can also include a target check valve **3978** positioned in the target fluid pathway to allow fluid to flow from the syringe **3912** to the IV bag and prevent fluid from flowing from the IV bag back toward the syringe **3912**. The source and target check valves **3976**, **3978** can be duck bill check valves similar to the check valve **2900** discussed herein, although dome check valves or disc check valves or any other suitable check valve can be used.

FIG. **54** is a cross sectional view of the connector **3910** and syringe **3912** showing fluid flowing through the connector **3910** from the vial **3907** to the syringe **3912**. As the plunger of the syringe **3912** is withdrawn, fluid is drawn into the syringe. The pressure causes the source check valve **3976** to open so that fluid is allowed to flow from the vial **3907** to the syringe **3912**. The pressure also causes the sides of the target check valve **3978** to bear against each other to maintain the target check valve **3978** closed. Thus, fluid drawn into the syringe **3912** will be drawn from the vial **3907** and not the IV bag. As fluid is drawn out of the vial **3907**, air can enter the vial **3907** through the air inlet channel **3946** as described above in connection with FIG. **43**.

FIG. **55** is a cross sectional view of the connector **3910** and syringe **3912** showing fluid flowing through the connector **3910** from the syringe **3912** toward the IV bag assembly **3914**. As the plunger of the syringe **3912** is advanced, fluid is driven out of the syringe. The pressure causes the target check valve **3978** to open so that fluid is allowed to flow from the syringe **3912** toward the IV bag assembly **3914**. The pressure also causes the sides of the source check valve **3976** to bear against each other to maintain the source check valve **3976** closed. Thus, fluid driven out the syringe **3912** will be directed to the IV bag and not back into the vial **3907**.

FIG. **56** is a perspective view of the IV bag assembly **3914**. The IV bag assembly **3914** can include an IV bag **3980**, a length of tubing **3982**, and a female connector **3984**. The female connector **3984** can be removably or irreversibly attached to the tubing **3982**. The female connector **3984** can function to seal off the IV bag assembly **3914** so that no fluid can escape from the IV bag **3980** except when a male connector is attached thereto.

FIG. **57** is an alternative IV bag assembly **5700** which may be used with the fluidics assembly **3906** or with various other embodiments discussed herein. The IV bag assembly **5700** can include an IV bag **5702** and a length of tubing attached thereto **5704**. A spike port **5706** can be positioned at the end of the tubing **5704**, and the spike port **5706** can include a

62

piercing membrane or barrier that when closed prevents fluid from entering or exiting the IV bag **5702**. The female connector **5708** can have a spike **5710** attached thereto. The spike **5710** can be inserted into the spike port **5706** until it pierces the membrane or barrier thereby providing access to the interior of the IV bag.

FIG. **58** is a perspective view of the top connector **3916** which includes a base member **4002** and a cassette **4004** in an engaged configuration. FIG. **59** is an exploded perspective view of the top connector **3916** with the base member and cassette **4004** in a disengaged configuration. FIG. **60** is a right-side view of the top connector **3916**. FIG. **61** is a front view of the top connector **3916**. FIG. **62** is a back view of the top connector **3916**. FIG. **63** is a left-side view of the top connector **3916**. FIG. **64** is a top-down view of the top connector **3916**. FIG. **65** is a bottom-up view of the top connector **3916**. FIG. **60** is a right-side view of the top connector **3916**. FIG. **60** is a right-side view of the top connector **3916**. FIG. **61** is a front view of the top connector **3916**. FIG. **62** is a back view of the top connector **3916**. FIG. **63** is a left-side view of the top connector **3916**. FIG. **64** is a top-down view of the top connector **3916**. FIG. **65** is a bottom-up view of the top connector **3916**.

FIG. **66** is a front view of the cassette **4004**. FIG. **67** is a back view of the cassette **4004**. FIG. **68** is a right-side view of the cassette, **4004**. FIG. **69** is a top-down view of the cassette **4004**. FIG. **70** is a bottom-up view of the cassette **4004**. FIG. **71** is a left-side view of the cassette **4004**.

FIG. **72** is a front view of the base member **4002**. FIG. **73** is a back view of the base member **4002**. FIG. **74** is a right-side view of the base member **4002**. FIG. **75** is a top-down view of the base member **4002**. FIG. **76** is a bottom-up view of the base member **4002**. FIG. **77** is a left-view of the base member **4002**.

The top connector **3916** can have features that are similar to, or the same as, the top connector **1900**, or any other suitable top connector discussed herein. For example, the top connector can include a light source and sensor to detect an air bubble in the connector **3910**, which can be an indication that the vial **3907** is empty. In some instances, infrared light can be used to detect the presence of air in the connector **3910**. For example, in some embodiments, light having a wavelength of at least about 980 nanometers and/or no more than about 1180 nanometers, or of at least about 1050 nanometers and/or no more than about 1110 nanometers, or of approximately 1080 nanometers can be effective for detecting air in the connector **3910**. Other wavelengths of light can also be used, such as light having a wavelength of at least about 850 nanometers and/or no more than about 1050 nanometers, or of at least about 920 nanometers and/or no more than about 980 nanometers, or of approximately 950 nanometers. Light can be used that has a wavelength of at least about 1380 nanometers and/or no more than about 1580 nanometers, at least about 1450 nanometers and/or no more than about 1510 nanometers, or about 1480 nanometers. One suitable optical sensor that can be used is the DL20JJ 1480 nm sensor available from STM Sensor Technologie Munchen GmbH of Germany. Light can be directed between hole **4006a** and hole **4006b** (hidden from view). The sensor region **3974** of the connector **3910** can be positioned between hole **4006a** and hole **4006b** when it is properly attached to the top connector **3916**.

In various embodiments disclosed herein which use a light source and a light sensor (e.g., to detect air or to detect the presence of an IV bag), the light source can pulse or flash at a predetermined frequency, and the light sensor can be configured to synchronize with the pulsing light source. In some embodiments, the light sensor can be configured to ignore

63

light that is not pulsed at the predetermined frequency. Thus, the light sensor can differentiate between light emitted by the corresponding light sensor (which is pulsed at the predetermined frequency) and light emitted from other sources (e.g., light from a different sensor that is pulsed at a different frequency, or ambient light). In some embodiments, light sources can be used that provide a constant beam of light.

The top connector **3916** can also include a light source and sensor configured to detect whether an IV bag assembly **3914** is attached to the connector **3910**. Light can be directed from hole **4008a** to hole **4008b** (hidden from view) and can intersect the second male connector **3968** at a location that is not obstructed when the second male connector **3968** is closed (when no IV bag is attached) and is obstructed when the second male connector **3968** is open (when an IV bag is attached). For example the location where the light intersects the second male connector **3968** can be the location **4012** shown in FIG. **78**. FIG. **78** is a cross sectional view of the second male connector **3968** in the closed configuration, with no IV bag assembly attached thereto. The light can pass through the clear housing **4016** unobstructed when the second male connector **3968** is in the open configuration. When the light reaches the corresponding detector, a signal can be generated that indicates that no IV bag is attached to the second male connector **3968**. When the valve member **4018** of the second male connector **3968** is pushed back to the open configuration (when the IV bag is attached), the opaque valve member **4018** is positioned to occupy the location **4012** and obstruct the light from reaching the corresponding detector. When no light reaches the detector, a signal can be generated that indicates that the second male connector **3968** is in the open configuration and the IV bag assembly **3914** is attached.

One suitable optical sensor that can be used with some embodiments for detecting the presence of IV bag or other target container is the DL20RM 645 nm sensor available from STM Sensor Technologie Munchen GmbH of Germany. In some embodiments, an amplifier can be used to amplify the signal of the light detector so that a relatively small amount of light can trigger the sensor. Thus, the amplifier can allow the sensor to accurately identify a closed valve member **4018** in the second male connector **3968** even when a portion of the light is reflected or refracted or otherwise redirected away from the light detector. One suitable amplifier that can be used is the V8-C or V8-D amplifier available from STM Sensor Technologie Munchen GmbH of Germany.

The top connector **3916** can also include a light source and detector configured to detect the presence of the second male connector **3968** regardless of whether it is open or closed. Light can be directed between hole **4010a** to hole **4010b** which is aligned with an opaque portion of the second male connector **3968**, e.g., at location **4014** as shown in FIG. **78**. When light passes unobstructed between hole **4010a** and hole **4010b** (hidden from view) the detector can generate a signal indicating that the connector **3910** (of which the second male connector **3968** is a part) is not present. When the light is obstructed by the plunger at location **4014** and does not reach the detector, a signal can be generated that indicates that the second male connector **3968**, and the rest of the connector **3910** is present.

In some embodiments, the two optical sensors can both function to detect whether an IV bag is attached. As further described below, if the light from one of the optical sensors is unintentionally blocked from reaching the corresponding light detector when the valve member is closed and no IV bag is present, the light from the other optical sensor can reach the corresponding light detector to provide an indication that the valve member is closed.

64

FIG. **79** is a perspective view showing the top connector **3916** cut to reveal the inner channels used to route wires for the light sources and detectors described above. FIG. **80** is a perspective view showing the top connector **3916** cut along a different axis to further reveal the channels used to route wires. Wires can pass from the main housing **3902** to the top connector **3916** via the hole **4020**. The wires can then enter the channel **4016** which leads to the holes **4006a-b**. As seen in FIG. **80**; the channels **4016** turn upward and lead to the holes **4008a-b** and the holes **4010a-b**.

In some embodiments, the cassette **4004** can be shaped or otherwise configured to be compatible with only authorized connectors **3910**. For example, as can best be seen in FIG. **61** (front view of the top connector **3916**), the side walls **4003** of the cassette **4004** are slanted. The slanted side walls can correspond to the slanted side walls of the lower housing portion **3962** of the connector **3910**. When an authorized connector **3910** specifically designed for use with the fluid transfer system **3900** is attached to the top connector **3916**, the tapered walls can fit snugly to properly position the connector **3910**. If an unauthorized connector of different size or shape were to be connected to the top connector, it would not fit properly with the top connector **3016**. The tapered walls can reliably position the connector **3910** with little or no freedom of movement in the vertical direction when the connector **3910** is attached to the top connector **3916**. The side walls can also restrict the freedom of movement of the connector along a horizontal direction that intersects the side walls.

It can be beneficial to limit the connectors that can be used with the system **3900** to ensure accurate and reliable transfer of fluid. For example, as discussed below, in some embodiments, the proper priming of the connector **3910** relies in part on the internal volume of the connector **3910**. Thus, if a different connector **3910** having a different internal volume were used, the system **3900** may improperly prime the connector **3910**.

In some embodiments, the top connector **3916** can be configured to hold the fluidics assembly **3906** in place using a securing mechanism. FIG. **81** is a perspective view of the base member **4002** of the top connector **3916** and the syringe **3912** cut and separated to reveal a channel **4022**. FIG. **82** is a top-down view taken at the cutting plane of FIG. **81**. The channel **4022** can be positioned such that when the syringe **3912** is fully attached to the top connector **3916**, the central axis of the syringe **3912** is positioned slightly past the central axis through the channel **4022**. As shown in FIG. **82**, one or more securing mechanisms **4024** can be positioned in the channel **4022**. In their relaxed position, the securing mechanisms **4024** can protrude partially past the channel **4022** and into the space shown occupied by the syringe **3912**. The securing mechanisms **4024** can be resiliently movable along the axis down the channel **4022**. As the syringe **3912** is slid into the top connector **3916**, the outer walls of the syringe **3912** contact the securing mechanisms **4024** and displace them into the channel **4022**. Once the widest portion of the syringe **3912** clears the securing mechanisms **4024**, the securing mechanisms **4024** return at least partially to their previous position, thereby securing the syringe **3912**, and the rest of the fluidics assembly **3906** in place. The securing mechanisms **4024** can attach the fluidics assembly **3906** to the top connector **3916** with little or no freedom of movement in the horizontal direction that is substantially perpendicular to the channel **4022**. By restricting the freedom of movement of the connector **3910**, the connector **3910** can reliably be aligned with respect to the optical sensors when it is attached to the top connector **3916**.

65

In some embodiments, the tray **3922** can be positioned as shown in FIG. **39** when in use and can be pivoted downward when not in use. The base member **4002** can be configured to facilitate the pivoting of the tray **3922**. FIG. **83** is a right-side view of the base member **4002** with the tray **3922** attached thereto. FIG. **84** is a right-side view of the base member **4002** and the tray **3922** in a disengaged configuration. The tray **3922** can have a rear connector **4026** and a front connector **4028**. The base member **4002** can include a rear connection slot **4030** that turns rearward and a front connection slot **4032** that turn forward. It will be understood that the other side of the tray **3922** and base member **4002** can be symmetrical or similarly configured. To attach the tray **3922** to the base member **4002**, the rear connector **4026** can be inserted into the rear connection slot **4030** until the rear connector **4026** reaches the rear depression **4034**. At this point the tray **3922** can hand from the top connector base member **4002** in the pivoted-down, unused position. The tray **3922** can be pivoted up until the forward connector **4028** enters the forward connection slot **4032**, and the tray can be shifted forward to the in-use position shown in FIG. **83** where the forward connector **4028** engages the forward depression **4036**.

In some embodiments, the system **3900** (or other systems described herein) can prime the fluidics assembly **3906** before the desired volume of fluid is transferred from the vial **3907** to the IV bag **3980**. When the user first assembles the fluidics assembly, the internal volumes contain air. FIG. **85** is flowchart that schematically shows an example embodiment of a method **8500** for priming a fluidics assembly.

At block **8504** a prime command is received. In some embodiments, the user can initiate the prime by providing an instruction to the system **3900** to prime the fluidics assembly. In some embodiments, the system **3900** can ask the user (via a user interface) whether the fluidics assembly should be primed. In some embodiments, the system can recognize when a new fluidics assembly has been attached to the system. For example the sensor that detects the presence of the second male connector can indicate when a fluids assembly was added to the system. Also, in some embodiments, other sensors can be used. The sensor for detecting air in the connector can also be configured to recognize whether the connector itself is present in the light path. Other sensor types are also possible. For example the securing mechanisms discussed above can include a sensor for detecting whether they are displaced, indicate that the connector is present. In some embodiments, the sensor that is used to detect air for determining whether vial has run empty can also be used to indicate whether the connector has already been primed by determining whether air is present in the connector. Thus, the system can be configured to determine when to automatically prime the fluidics assembly and when to prompt the user to decide whether to prime.

At block **8506** the method determines whether the fluidics assembly is properly attached. For example, the sensors discussed above can be used to determine whether the fluidics assembly is present and whether a prime is needed. In some embodiments, this step is performed before block **8504**, as discussed above. If the fluidics assembly is not properly attached, block **8508** can inform the user to attach or correct the fluidics assembly. If the fluidics assembly is properly attached, the method **8500** advances to block **8510**.

At block **8510**, the syringe plunger is withdrawn by the distance necessary to draw the priming volume into the syringe. The system can ignore the signal from the air detector when priming the fluidics assembly. Normally, the air detector can be used to prevent air from being drawn into the

66

syringe. However, during the priming process, air can be drawn into the syringe before the fluid reaches the syringe.

In some embodiments, the priming volume is the volume of the fluidics assembly between (and excluding) the vial and the IV bag assembly when the syringe plunger is fully advanced. The priming volume can be the volume of air in the fluidics assembly that needs to be pushed into the IV bag in order to bring the leading edge of fluid up to the entrance to the IV bag, which may be the end of a connector attached to the bag via a length of tubing. Thus, using the system **3900** as an example, the priming volume can, for example, be equal to the internal volume of the vial adapter **3908**, plus the internal volume of the connector **3910** (which includes the internal volume of the both male connectors **3964**, **3968**, the internal volume in the internal chamber with the check valves, and the internal volume of the syringe interface that is not occupied by the syringe). In some embodiments, the internal volume of the IV bag assembly is excluded from the priming volume. However, in some embodiments the internal volume of the female connector **3984** and the tubing **3982** and any other portions of the IV bag assembly other than the IV bag itself are included. This can be useful if the parts of the IV bag assembly need to be replaced or removed prior to patient delivery. In some embodiments, the priming volume can include a portion of the syringe's internal volume, such as the internal volume of the syringe tip above the plunger's end. In some embodiments, the vial adapter can be self priming, in which case, the internal volume of the vial adapter can be excluded from the priming volume. For example, in some embodiments, the air in the fluid pathway of the vial can rise up into the vial such that the fluid from the vial advances to the end of the female connector of the vial adapter.

In some embodiments, the system **3900** can calculate the priming volume based on information acquired from the user or from sensors or otherwise. For example, the priming volume may vary depending on the model of vial adapter that is used or the model of syringe being used. The system **3900** can prompt the user for information to be used for calculating the priming volume. In some embodiments, the priming volume can be a predetermined amount. For example, the priming volume can about 0.7 milliliters.

At **8512** the system determines whether the IV bag is attached, for example. If the IV bag is not attached properly, the system prompts the user to properly attached the IV bag at **8514**. If the IV bag is attached, the method **8500** advances to Block **8516**. At **8516**, the syringe drive the priming volume into the connector, through the second male connector, and into the IV bag assembly. In some embodiments, the priming volume that is drawn into and expelled from the syringe contains both air and fluid. If calculated and executed properly, in some embodiments, the leading edge of the fluid from the vial will be positioned at the entrance to the IV bag assembly, or in some cases at the entrance to the IV bag itself. At block **8518** the method can optionally prompt the user that the fluidics assembly was successfully primed.

The method **8500** can be varied in many ways. For example, the checks at blocks **8506** and **8512** can be omitted or performed together or performed before block **8504**. In some embodiments, the system does not perform a separate priming procedure. Instead the system can merely add the priming volume to the first volume of fluid that is transferred through the fluidics assembly.

FIG. **86** is a flowchart schematically showing a sample embodiment of a method **8600** for transferring fluid from a vial to an IV bag. This method can be similar in some ways to the method **2400** discussed above. At block **8602**, the amount of fluid to be transferred is determined. At block **8604**, the

67

system determines whether the amount remaining to be transferred is greater than the maximum volume that can be transferred by the syringe. If that remaining volume to be transferred is larger than the maximum volume of the syringe, the method proceeds to block **8606** where the system fills the syringe with the maximum syringe fluid volume. As fluid is drawn into the syringe, the air detector monitors for the presence of air in the connector, as will be discussed in greater detail in connection with FIG. **87**.

At block **8608**, the fluid is transferred from the syringe into the IV bag. In some embodiments the system can first perform a check to ensure that the IV bag is properly attached before advancing the plunger of the syringe. At block **8610**, the maximum volume of the syringe is subtracted from the volume to be transferred, and the process returns to Block **8604**.

Once the amount of volume to be transferred is less than the maximum volume of the syringe, the process advances to block **8612** where the system fills the syringe with the remaining amount of volume to be transferred. Again, while the fluid is drawn into the syringe, the air detector monitors for the presence of air in the connector, as will be discussed in greater detail in connection with FIG. **87**. At block **8614** the fluid is driven from the syringe into the IV bag. In some embodiments, the system can perform a check to ensure that the IV bag is properly attached before pushing fluid into the IV bag. The process then ends at block **8616**.

FIG. **87** is a flowchart that schematically illustrates an example embodiment of a method for replacing a vial of fluid to be transferred. At block **8702**, the air detector identifies air in the connector, and at block **8704** the system stops the transfer of fluid. In some embodiments, the system can prompt the user that air was detected and ask the user to check the vial. In some embodiments, the user interface can allow the user to indicate that the vial is not yet empty, in which case, the detected air was likely merely a small bubble. If the system receives notification that the vial is not empty at block **8706**, the process will then continue transferring the fluid at block **8708**.

If the vial was indeed empty, the user can replace the vial and the corresponding vial adapter. In some embodiments, the user can press a button or otherwise indicate that the vial has been replaced. Once notification is received that the vial has been replaced at block **8712**, the system then adds a replacement volume amount to the target fluid transfer amount to compensate for the volume of air that was drawn from the vial before the air was detected. In some embodiments, the vial replacement volume can be substantially equal to the internal volume of the flow path through the vial adapter, through the first male connector, and through the portion of the connector that is on the syringe side of the target check valve and before the sensing location where the air was detected. In some embodiments, the volume of the flow path through the new vial adapter should also be added to the vial replacement volume since the air in the new vial adapter will also be drawn into the syringe and then pushed to the IV bag. As discussed above, variations are possible. For example, for a self priming vial adapter, the volume for the replacement vial adapter does not need to be included. In some embodiments, the vial replacement volume can be 0.3 milliliters.

At block **8716** the method continues with the fluid transfer process. In some embodiments, the system can ignore air detected in the connector for a short time after the vial is replaced. In some embodiments, after the vial replacement volume has been added to the total transfer volume, the system can reevaluate whether an additional syringe draw will be needed to reach the desired total fluid transfer amount.

68

FIG. **88** is a perspective view of another example embodiment of a fluid transfer system **8800**. The fluid transfer station **8800** can be similar to, or the same as, fluid transfer systems **3900**, **100**, or **600** or any other fluid transfer system discussed herein. Thus, the discussion associated with many features of other fluid transfer systems described herein is also applicable to the fluid transfer system **8800**, even when not specifically identified.

The fluid transfer system **8800** can include a main housing **8802** that supports four fluid transfer stations **8804a-d**, although any other suitable number of fluid transfer stations can be used. In the illustrated embodiment, the fluid transfer stations **8804a-b** are configured to receive larger syringes than the fluid transfer stations **8804c-d**. For example, fluid transfer stations **8804a-b** can be configured to use 20 milliliter syringes and fluid transfer stations **8804c-d** can be configured to use 10 milliliter syringes, although other sizes of syringes can also be used. In some embodiments, a larger syringe (e.g., 20 milliliters) can allow fluid to be transferred from the source container to the target container at a faster rate, while a smaller syringe (e.g., 10 milliliters) can allow fluid to be transferred from the source container to the target container with greater precision. It will be understood that the fluid transfer stations **8804a-d** can be configured to use various other syringe sizes, such as syringes of sizes between about 1 milliliter and about 100 milliliters or even syringes outside these ranges.

The fluid transfer station **8804d** is shown as having a fluidics assembly **8806** attached thereto. The fluidics assembly can include a vial (not shown in FIG. **88**), a vial adapter **8808**, a connector **8810**, a syringe **8812**, and an IV bag assembly **8814** (partially shown in FIG. **88**), which can be similar to, or the same as, the corresponding components discussed in connection with the embodiment shown in FIG. **39**, or any other embodiments disclosed herein. The transfer station **8804d** can be configured to receive the syringe **8812** and/or the connector **8810** using, for example, a top connector **8816**, a middle connector **8818**, and a lower connector end piece **8820**. A motor (hidden from view in FIG. **88**) can cause the lower connector **8820** to move to withdraw and advance the plunger of the syringe **8812**. As discussed above, the motor can be a high precision stepping motor.

The fluid transfer station **8804d** can include a tray **8822** to support the IV bag (not shown in FIG. **88**). The tray **8822** can be attached to the top connector **8816** by a tray arm **8824** as will be discussed in greater detail below. The housing **8802** can include a step or foot **8830** positioned at the base thereof to provide increased stability to the housing **8802**, for example to prevent the weight of the IV bags from tipping the housing **8802** forward.

FIG. **89** is a perspective view of the top connector piece **8816**. The top connector piece can be similar to, or the same as the top connector pieces **3916** or **1900** or any other top connector piece described herein. The top connector **8816** can include a base member **8902** and a removable cassette **8904**. The base member **8902** can include a tray hole **8906** that is configured to receive the tray arm **8824** therein. The tray hole **8906** can be positioned near a side edge of the base member **8902** and the tray arm **8824** can similarly be attached near a side edge of the tray **8822** (as seen in FIG. **90**). Thus, the tray **8822** can be positioned substantially centered in front of the top connector **8816** while the tray arm **8824** is offset to the side so that the tray arm **8824** does not interfere with the attaching and detaching of the IV bag assembly.

With further reference to FIG. **90**, the tray arm can have a substantially circular cross-sectional shape, or can otherwise be configured to allow the tray arm **8824** to rotate within the

69

tray hole 8906. The tray arm 8824 can include a notch 8826 formed in the end opposite the tray 8822. The tray arm 8824 can also include a groove 8828 that extends around all or part of the circumference of the tray arm 8824.

FIG. 91 shows a rear perspective view of the top connector 8816 with the tray 8822 attached thereto in a first configuration wherein the tray 8822 is positioned to support an IV bag. FIG. 92 shows another rear perspective view of the top connector 8816 with the tray 8822 attached thereto in a second configuration wherein the tray 8822 is pivoted by about 90° to provide unobstructed access to the cassette 8904. The user can, for example, pivot the tray 8822 out of the way to the second configuration (shown in FIG. 92) when attaching the syringe 8812 and/or the connector 8810 to the fluid transfer station 8804d. Then the user can pivot the tray 8822 back to the first configuration (shown in FIG. 91) and place the IV bag onto the tray 8822.

The top connector 8816 can include a stop plate 8908, which can be positioned to occupy a portion of the tray hole 8906. The stop plate 8908 can be secured to the back surface of the base member 8902 using, for example, a screw 8910, and the back surface of the base member 8902 can have a recess shaped to receive the stop plate 8908 therein. The stop plate 8908 can have a thickness that is configured to fit into the notch 8826. When the tray 8822 is in the first configuration (shown in FIG. 91), the wall of the notch 8826 abuts against the side surface of the stop plate 8908 to prevent the tray 8822 from rotating past the first configuration. When the tray 8822 is rotated to the second configuration (shown in FIG. 92), the wall of the notch 8826 abuts against the bottom surface of the stop plate 8908 to prevent the tray 8822 from pivoting past the second configuration. In the illustrated embodiment, the stop plate 8908 is generally square shaped, such that the tray 8822 pivots by at least about 75° and/or no more than about 105°, or in some cases about 90° between the first configuration and the second configuration. The shape of the stop plate 8908 and/or the shape of the notch 8826 can be modified to change the rotational distance between the first and second tray configurations. For example, in some embodiments, the tray can pivot by about 180°, or by any angular distance, between the first and second configurations. Also, the notch and/or the stop 8826 plate 8908 can be moved or modified so that the tray 8822 rotates in the opposite direction of that shown in FIGS. 91-92.

FIG. 93 is a perspective view of the top connector 8816 and the tray arm 8824 cut along a vertical plane that intersects the axis of the tray hole 8906. A top hole 8912 can be formed in the base member 8902 and can intersect the tray hole 8906. When the tray arm 8824 is inserted into the tray hole 8906, the groove 8826 can align with the top hole 8912. A securing mechanism 8914 can be positioned in the top hole 8912 so that the securing mechanism 8914 can interface with the groove 8826 to secure the tray arm 8824 into the tray hole 8906. The securing mechanism 8914 can have a tip 8916 that is attached to a spring such that the tip 8916 can be axially displaced along the top hole 8912 in a direction away from the tray hole 8906 to compress the spring. When the tray arm 8824 is inserted into the tray hole 8906, the tray arm 8824 displaces the tip 8916 of the securing mechanism 8914 and compresses the spring. Once the tray arm 8824 is inserted far enough for the groove 8826 to align with the securing mechanism 8914, the tip 8916 can snap down into the groove 8826. Thus, the securing mechanism 8914 can prevent the tray arm 8824 from being accidentally removed from the tray hole 8806. To remove the tray arm 8824 from the tray hole 8906, the user can pull the tray arm 8824 with enough force to

70

compress the spring the drive the tip 8916 out of the groove 8826. The groove 8826 can be V-shaped to facilitate the removal of the tray arm 8824.

Although not shown in the illustrated embodiment, the groove 8826 can include deepened portions that are configured to receive the tip 8916 when the tray 8822 is in the first configuration and in the second configuration, so that the tray 8822 can be “locked” into the first configuration or into the second configuration. To break the “lock” and allow the tray 8822 to pivot, the user can apply a rotational force that is sufficient to compress the spring and drive the tip 8916 out of the deepened portion of the groove 8826. In some embodiments, the groove 8826 can be omitted, and the tray arm 8824 can include two holes configured to receive the tip 8916 when in one of the first and second configurations.

With further reference to FIG. 93, a cap 8918 can be placed over the top opening of the top hole 8912 to prevent debris from entering the hole 8912. Two bushings 8920, 8922 can be positioned in the arm hole 8906, one near the stop plate 8908, and the other near the opening of the arm hole 8906. Other numbers of bushings can be used, or the bushings can be omitted. The bushings 8920, 8922 can be made from a compressible material and can have openings that are slightly smaller than the diameter of the tray arm 8824. Thus, the tray arm 8824 can compress the bushings 8920, 8922 as the tray arm 8824 is inserted into the tray hole 8906. The pressure applied to the tray arm 8824 by the bushings 8920, 8922 can provide additional stability to the tray 8824 to prevent rattling or accidental rotation.

FIG. 94 is a cross sectional view of the top connector 8816 and tray arm 8824 taken along a horizontal plane that intersects the axis of the tray hole 8906. A channel 8824 can extend through the base member 8902, and securing mechanisms 8926, 8928 can be positioned in the channel 8924 so that the tips 8930, 8932 thereof extend out from the channel 8824. In the illustrated embodiment, the channel 8924 can intersect the tray hole 8906. As similarly discussed in connection with FIG. 82, when a syringe is attached to the top connector 8816, the syringe can displace the tips 8930, 8932 into the channel 8824 to compress the springs of the securing mechanisms 8826, 8828. Once the widest portion of the syringe passes the tips 8930, 8932, the springs can drive the tips 8930, 8932 toward each other to secure the syringe to the top connector 8816. Securing mechanisms can similarly be used to secure other portions of the fluidics assembly 8806 (e.g., the connector 8810, or vial adapter 8808) to the transfer station 8804d.

FIG. 95 is a perspective view of the cassette 8904, which can be similar to, or the same as, the cassette 4004, 1904, or any other suitable cassette described herein. The cassette 8904 can include holes 8940a-b that are configured to provide light path between a light source and a light sensor configured to detect air in the connector 8810. The cassette 8904 can also provide holes 8942a-b and holes 8944a-b to provide light paths between corresponding light sources and light detectors for detecting the presence of an IV bag assembly. The cassette 8904 can include channels 8946 configured to provide a path for wires to reach the light sources and light detectors. The wires can pass through a hole in the base member 8902 (not shown in FIG. 95) and through a hole 8948 that leads to the channels 8946. One channel can lead to the holes 8942b and 8944b used in detecting the presence of the IV bag, and another channel can branch off and lead to the hole 8940b used for detecting air. The other side of the cassette 8904 can have similar channels leading to the holes 8904a, 8942a, and 8944a. As discussed herein, the cassette 8904 can be removably attachable (e.g., using a screw) to the base member 8902,

71

so that the cassette **8904** can be detached to provide access to the channels **8946** and to the light sources and light detector, if, for example, a component needs to be repaired or replaced.

The cassette **8904** can have side walls **8950** that are tapered similar to the cassette **4004** disclosed above. In the illustrated embodiment, the cassette **8904** has vertical side walls **8950** that are not tapered (as can be seen in FIG. 96).

FIG. 97 is a cross sectional view of the connector **8810** with an outline of the cassette **8904** shown in dotted lines. In the illustrated embodiment, the hole **8940a** for the air sensor aligns with the fluid pathway through the transition between the source connector piece **8952** and the main connector body **8954**. Thus, the light used to detect air passes through a wall of the female end **8956**, through a wall of the male end **8958**, through the fluid pathway **8960**, then through an opposite wall of the male end **8958**, and through an opposite wall of the female end **8956**. At least a portion of the female end **8956** and at least a portion of the male end **8958** can be substantially transparent to the light used for the air sensor. In some cases, at least the entire pieces that are integrally formed with the female end **8956** and the male end **8958** can be substantially transparent to the light of the air sensor.

The air detection light can intersect the fluid pathway at a location of the fluid pathway between the source check valve **8962** and the source container (not visible in FIG. 97). In some cases, detecting air bubbles at a location upstream from the source check valve **8962** can reduce the occurrence of false air bubble reads which can result from the turbulent flow of fluid through the source check valve **8962** even when the source container has not run dry. In some embodiments, the light for the air sensor can pass through a fluid passageway that is less than about 4 millimeters wide, or less than about 2 millimeters wide; and the fluid passageway can be less than about quadruple the size, less than about triple the size, less than about double the size, or no larger than the hole **8940a** associated with the light for the air sensor. By causing the light from the air sensor to cover a large portion of the fluid pathway, the sensor can more reliably identify the leading edge of air when the source container has run dry.

FIG. 98 is a perspective view of a connector **9800** which can be similar to the connector **8810**, or any other connector disclosed herein. A male end **9806** of the source connector piece **9804** can connect to a female end **9808** of the main body portion **9802** of the connector **9800**. The female end **9808** can have substantially flat outer surfaces **9810** where the light from the air sensor intersects the female end **9808** to enter the connector **9800**, so that the light enters the connector at a direction that is substantially normal to the surface **9810** (e.g., within about 10° or 5° or less of a direction normal to the surface **9810**), thereby reducing the likelihood that the light will be refracted, or otherwise misdirected, away from the light sensor.

In the embodiment illustrated in FIG. 98, the inner surface of the female end **9808** is curved and tapered so as to receive the curved and tapered outer surface of the male end **9806**. However, in some embodiments, additional surfaces that intersect the light from the air sensor can be flat. For example, at least a portion of the outer surfaces and the inner surfaces of the male end **9806** and at least a portion of the inner surfaces of the female end can also be flat. In some embodiments, each surface that the light for the air sensor passes through on the female end **9810** and the male end **9806** is a flat surface. In some embodiments, the male end **9806** and the female end **9808** can be substantially index matched when they are mated together, thereby reducing refraction, or other misdirection, of the light away from the corresponding sensor.

72

Returning now to FIG. 97, the target connector piece **8964** can align with the holes **8942a** and **8944a** which are associated with two optical sensors used for detecting an IV bag. In the illustrated embodiment, two optical sensors can be used to determine whether an IV bag is attached to the target connector piece **8964**. As shown in FIG. 97 by the positions of the holes **8942a** and **8944a**, a first light path can pass through the target connector piece **8964** at a location above the outside surface of the plunger **8966**, and a second light path can pass through the side wall of the plunger **8966**. As similarly explained in connection with FIG. 19D, when no IV bag is attached to the target connector **8964**, the valve member **8970** can be positioned in an open position, as shown in FIG. 97, to allow light to pass through the transparent components of the target connector piece **8964** to the corresponding light detectors. When the light detectors detect the light, they can provide a signal indicating that the no target container is attached to the target connector piece **8964**. In response to that signal, a controller can stop or prevent the transfer of fluid thereby preventing fluid (e.g., hazardous chemotherapy drugs) from being sprayed out of the target connector piece **8964** when no IV bag is attached thereto. In a manner similar to that discussed in connection with FIG. 19E, when a connector of an IV bag assembly is attached to the target connector piece **8964**, the valve member **8970** can be displaced to an open position in which an opaque portion of the valve member **8970** is positioned in between the holes **8942a** and **8942b** and also between the holes **8944a** and **8944b**, to block light of the optical sensors from reaching the light detectors. When the light detectors do not detect the light, they can provide a signal indicating that a target container is attached to the target connector piece **8964**. In response to the signal, a controller can begin, resume, or allow the transfer of fluid through the connector.

In some embodiments, the connector **8810** can attach to the transfer station with some freedom of movement. Thus, in some instances, the light paths may not align at the precise locations shown. In some instances, one of the light paths may intersect the fluid pathway **8968** through the plunger **8966**. Accordingly, a frequency of light can be used that is not blocked by the fluid (e.g., chemotherapy drugs) being transferred through the connector **8810**. In some embodiments, a wavelength of light can be used that transmits well through water or saline, which can be used as a solvent or diluent for the drugs. In some embodiments, visible light can be used (e.g., red colored light). In some embodiments, light can be used for IV bag detection that has a wavelength of at least about 545 nanometers and/or no more than about 745 nanometers, or of at least about 615 nanometers and/or no more than about 675 nanometers, or of about 645 nanometers.

The embodiment of FIG. 97 includes two optical sensors for detecting an IV bag, and the controller can be configured to only allow fluid to be transferred through the target connector piece **8964** when both of the light detectors do not detect light from their corresponding light sources. While no IV bag is attached, if light from one of the optical sensors is unintentionally blocked or diverted away from the corresponding light detector, the light from the other optical sensor can reach its corresponding light detector, thereby preventing a false read in which the controller receives a signal that an IV bag is attached when no IV bag is present. Light from one of the optical sensors can be unintentionally blocked or diverted by various different causes.

As mentioned above, in some cases, the connector **8810** can connect to the fluid transfer station with some freedom of movement. Thus, in some instances, one of the light beams from one of the optical sensors may strike the curved housing

73

8972 of the target connector piece **8964** at a location other than at the locations shown in FIG. **97** associated with the holes **8942a** and **8944a**. If the connector is shifted enough from the position shown in FIG. **97**, one of the light beams can strike the curved housing **8972** at a sufficiently oblique angle so that the light is reflected, refracted, or otherwise unintentionally diverted from its normal substantially linear path through the target connector piece **8964**. Thus, the light can fail to reach the corresponding light detector even when the valve member **8970** is in the closed position.

The light path formed between the holes **8942a** and **8942b** can be spaced from the light path formed between the holes **8944a** and **8944b** in a direction transverse to the longitudinal axis of the target connector portion. The distance can be sufficient so that if one of the light paths intersects the curved housing **8972** at an angle that is oblique enough to divert the light, the other light path will travel through the target connector piece **8964** at a location close enough to the longitudinal axis so that the light strikes the curved housing **8972** at an angle that is close enough to normal so that the light is not diverted away from the corresponding light detector. For example, the holes **8944a** and **8944b** can be positioned substantially directly below the holes **8942a** and **8942b**. The hole **8944a** can be spaced away from the hole **8942a** by a distance of at least about 2 millimeters and/or no more than about 6 millimeters, or by about 4 millimeters. The hole **8944b** can be spaced away from the hole **8942b** by substantially the same distance.

As similarly discussed above, in some embodiments, the connector **8810** can be secured to the top connector **8816** such that it has little or no freedom of movement so that the connector **8810** can reliably be aligned with the optical sensors.

FIGS. **99-104** are cross sectional views of the target connector piece **8964** taken along the line **99-99** in FIG. **97**. FIGS. **99-104** show how different rotational positions for the housing **8972** can affect the light of the two optical sensors. As previously discussed, the housing **8972** of the target connector piece **8964** can have gaps **8974a-b** formed therein. In some embodiments, the light of one of the optical sensors can be scattered, reflected, refracted, or otherwise unintentionally blocked from reaching the corresponding light detector when an edge of one of the gaps **8974a-b** is positioned between the light source and light detector. For example, the edges of the housing **8972** at the gaps **8974a-b** can have a generally rough surface that scatters light so that the edges are substantially opaque to the light from the optical sensors.

The optical sensors and the corresponding holes **8942a-b** and **8944a-b** can be positioned such that if one light path is obstructed by one of the gaps **8974a-b**, the other light path will not be obstructed. For example, in some embodiments, the light paths can be spaced from the center of the target connector piece **8964** by different amounts. For example, a first light path can be spaced about 3 millimeters from the center of the target connector piece **8964** and a second light path can be spaced about 1 millimeter from the center of the target connector piece **8964** in the opposite direction. Other orientations are also possible.

When the housing **8972** is oriented as shown in FIG. **99**, the light from the first light source **8976a** can travel through the target connector piece **8964** to the first light detector **8978a** without obstruction. Similarly, light from the second light source **8976b** can travel through the target connector piece **8964** to the second light detector **8978b** without obstruction. It will be understood that although the light can refract as it passes through certain surfaces of the target connector piece **8964**, the light can follow a substantially linear pathway

74

between the light sources **8976a-b** and the corresponding light detectors **8978a-b**, as shown by the dotted lines in FIG. **99**.

If the housing **8972** is rotated to the position shown in FIG. **100**, the light from the first light source **8976a** can strike an edge of the gap **8974b** and be blocked from reaching the first light detector **8978a**. However, the light from the second light source **8976b** can pass through the target connector piece **8964** to the second light detector unobstructed.

If the housing **8972** is further rotated to the position shown in FIG. **101**, the light from the second light source **8976b** can be obstructed by an edge of the gap **8974a**. However, in this orientation, the light from the first light source **8976a** can pass through the gap **8974b** without being obstructed by the edges thereof.

If the housing **8972** is further rotated to the position shown in FIG. **102**, the light from the first light source **8976a** can be obstructed by an edge of the gap **8974b**. However, the light from the second light source **8976b** can pass through the gap **8974a** without being obstructed by the edges thereof.

If the housing **8972** is further rotated to the position shown in FIG. **103**, the light from the second light source **8976b** is obstructed by an edge of the gap **8974a**. However, the light from the first light source **8976a** can pass through the target connector piece **8964** to the first light detector **8978a** without being obstructed, as shown.

If the housing **8972** is further rotated to the position shown in FIG. **104**, the light from both light sources **8976a-b** can pass through the target connector portion **8964** to the corresponding light detectors **8978a-b**, as shown.

In some embodiments, the target connector portion can be configured to be used with a single optical sensor for detecting whether the valve member is open or closed. For example, the target connector portion can be modified so that the gaps between the walls of the housing do not intersect the light path of the optical sensor.

FIG. **105** is a side view of another example embodiment of a connector **9000** which can be similar to, or the same as, the connector **8810**, the connector **3910**, the connector **320**, or any other suitable connector discussed herein. The connector **9000** can include a main body portion **9002**, a source connector portion **9004**, and a target connector portion **9006**, which can be similar to, or the same as, the corresponding components in, for example, the connector **8810**, the connector **3910**, or the connector **320**. The target connector portion **9006** can be similar to the target connector portion **338** discussed above, and much of the disclosure relating to the target connector portion **338** also applies to the target connector portion **9006**. FIG. **106** is a cross sectional view of the target connector portion **9006**.

With further reference to FIGS. **105** and **106**, the target connector portion **9006** can include a housing **9008**, a sealing ring **9009**, a valve member **9010**, a resilient member **9012**, a first end cap member **9014**, and a second end cap member **9016**. The sealing ring **9009**, valve member **9010**, resilient member **9012**, and second end cap member **9016** can be the same as the corresponding components of the target connector portion **338**. The first end cap member **9014** can be a modified version of the first end cap member **405** of the target connector portion **338**. The first end cap member **9014** can have forward wall portion **9022** that surrounds a portion of the plunger **9024** on the second end cap member **9016** when assembled. The housing **9006** can include a first wall **9018a** and a second wall **9018b** with gaps **9020a-b** formed therebetween to accommodate the elongate elastic members of the resilient member **9012**.

The housing **9006** can attach to the ends of the forward wall portion **9022** by sonic welding, adhesive, mechanical attachments, or any other suitable manner. The target connector portion can be attached to a corresponding fluid transfer station that includes one or more optical sensors so that the light path of the optical sensor passes through the forward wall portion **9022**. The first end cap member **9014** can be substantially transparent, and in some cases, the second end cap member **9016** can be substantially transparent as well. For example, the light path can pass through the target connector portion **9006** at a location within the area **9026** shown in dotted lines in FIG. **106**. In some cases, the light path can pass through the target connector portion **9006** at about the centerline through the connector (e.g., at location **9028**) such that the light enters and exits the curved surfaces of the forward wall portion **9022** at a direction that is substantially normal to the surfaces, thereby reducing the occurrence of unintentional redirecting of the light. Because the housing **9008** does not extend back into the light path, the gaps **9020a-b** in the housing **9008** do not obstruct the light. The forward wall portion **9022** can be an unbroken, generally cylindrical wall, at least in the area that intersects the light path of the optical sensor. Thus, a single optical sensor can be used to determine whether the valve member **9010** is in the open or closed configuration.

Many different connector types can be used for the source connector portion and/or the target connector portion of the various connectors disclosed herein. Various other connector types can include a valve member, or other movable component, that can be transitioned in and out of the light path of an optical sensor to indicate whether an IV bag is attached to the connector. FIG. **107** is a perspective view of an example embodiment of a connector **9100**. The connector **9100** can include a main body portion **9102**, a source connector portion **9104**, and a target connector portion **9106**. The connector **9100** that can be similar to the connector **3910** or **8810** except that the target connector portion **9106** can be a version of the Clave® connector manufactured by ICU Medical, Inc., of San Clemente, Calif. Various embodiments of a connector of this type are described in the '866 Patent. Additional details and alternatives are also provided in U.S. Provisional Patent Application No. 61/345,554, filed May 17, 2010, the entirety of which is hereby incorporated by reference herein.

The target connector portion **9106** can include a valve member **9108** disposed therein, which can transition between a closed position when no IV bag is attached thereto and an open position when an IV bag is attached thereto.

FIG. **108** is a cross sectional view of the target connector portion **9106** with the valve member **9108** in the closed configuration. FIG. **109** is a cross sectional view of the target connector portion **9106** with the valve member **9108** in the open configuration.

A housing member **9110** can attach to a base **9112** to define an interior chamber **9114** therein. The base can have a spike **9116** extending into the interior chamber **9114** and a male end **9118** extending generally opposite the spike **9116**. A fluid pathway **9120** can run through the spike **9116** and male end **9118**. The valve member **9108** can have a head **9122** that includes a slit **9124** therein. A resiliently compressible valve body **9126** can include a series of accordion sections or O-rings to bias the valve member **9108** toward the closed position. The end of the housing **9110** can be a female luer **9130** configured to receive a male luer end **9132** associated with, for example, an IV bag assembly.

In some embodiments, the housing member **9110**, or at least a portion thereof, can be substantially transparent, and the valve member, or at least a portion thereof, can be sub-

stantially opaque. Light from an optical sensor can pass through the housing **9110** and the interior chamber **9114** at a location **9128**. When the valve member **9108** is in the closed configuration, the light can travel through the target connector portion **9106** substantially unobstructed, to provide a signal indicating that the valve member **9108** is closed and no target container is attached. When the valve member **9108** is in the open configuration, it can be positioned in the light path such that the light is blocked from reaching the light detector. The light detector can then provide a signal indicating that the valve member **9108** is in the open configuration and a target container is attached thereto.

In some embodiments, the target connector portion can include an interaction portion. For example, in some embodiments, the interaction portion can comprise a generally opaque outer housing or can comprise a generally transparent outer housing and an internal generally opaque moveable portion. The optical sensor can be configured such that light is obstructed when the valve member is in the closed configuration and the light is permitted to pass to the light detector substantially unobstructed when the valve member is in the open configuration. For example, FIG. **110** is a cross sectional view of the target connector portion **9106** with the light path of the optical sensor passing through the target connector portion **9106** at a location **9134** that is blocked by the valve member **9108** when the valve member **9108** is closed (as shown in FIG. **110**) and is substantially unobstructed when the valve member **9108** is open (as shown in FIG. **111**). Accordingly, the controller can be configured to allow fluid transfer when the light detector is able to detect light transmitted through the target connector portion **9106** indicating that a source container is present, and the controller does not allow fluid transfer when the light detector does not detect the light.

It will be understood that various other types of connectors can be used for the target connector portion **9106** and can have a location where a light path is obstructed when the connector is in a first state (e.g., open or closed) and the light path is substantially unobstructed when the connector is in a second state (e.g., closed or open). Other variations are possible. In some embodiments, the optical sensor can be positioned to align with the connector of the IV bag assembly, or some other opaque portion of the IV bag assembly, such that when the IV bag assembly is present, the light is blocked from reaching the light detector to thereby generate a signal to allow fluid transfer.

Although many features of the embodiments shown in the Figures are specifically called out and described, it will be understood that additional features, dimensions, proportions, relational positions of elements, etc. shown in the drawings are intended to make up a part of this disclosure even when not specifically called out or described. Although forming part of the disclosure, it will also be understood that the specific dimensions, proportions, relational positions of elements, etc. can be varied from those shown in the illustrated embodiments.

Embodiments have been described in connection with the accompanying drawings. However, it should be understood that the foregoing embodiments have been described at a level of detail to allow one of ordinary skill in the art to make and use the devices, systems, etc. described herein. A wide variety of variation is possible. Components, elements, and/or steps may be altered, added, removed, or rearranged. Additionally, processing steps may be added, removed, or reordered. While certain embodiments have been explicitly described, other embodiments will also be apparent to those of ordinary skill in the art based on this disclosure.

Some aspects of the systems and methods described herein can advantageously be implemented using, for example, computer software, hardware, firmware, or any combination of software, hardware, and firmware. Software can comprise computer executable code for performing the functions described herein. In some embodiments, computer-executable code is executed by one or more general purpose computers. However, a skilled artisan will appreciate, in light of this disclosure, that any module that can be implemented using software to be executed on a general purpose computer can also be implemented using a different combination of hardware, software, or firmware. For example, such a module can be implemented completely in hardware using a combination of integrated circuits. Alternatively or additionally, such a module can be implemented completely or partially using specialized computers designed to perform the particular functions described herein rather than by general purpose computers.

While certain embodiments have been explicitly described, other embodiments will become apparent to those of ordinary skill in the art based on this disclosure. Therefore, the scope of the invention is intended to be defined by reference to the claims as ultimately published in one or more publications or issued in one or more patents and not simply with regard to the explicitly described embodiments.

The following is claimed:

1. A method of providing a fluid transfer module for use with a substantially entirely closed system for the transfer of medical fluids between or among different medical fluid containers by an electronically controlled fluid dispensing system, the method comprising:

(a) providing a fluid transfer module configured to be removably attachable to an electronically controlled fluid dispensing system, the fluid transfer module comprising:

(i) a first interface configured to be connected in fluid communication with a fluid source container, the first interface comprising a first selectively openable and closeable aperture that is configured to substantially entirely prevent fluid within the fluid transfer module from escaping through the first aperture when the first aperture is closed;

(ii) a second interface configured to be connected in fluid communication with a fluid destination container, the second interface comprising a second selectively openable and closeable aperture that is configured to substantially entirely prevent fluid within the fluid transfer module from escaping through the second aperture when the second aperture is closed;

(iii) an intermediate container, or an intermediate interface configured to be connected to an intermediate container;

(iv) a first valve configured to permit the passage of fluid from the fluid source container to the intermediate container and configured to generally obstruct the passage of fluid from the intermediate container to the fluid source container; and

(v) a second valve configured to permit the passage of fluid from the intermediate container to the fluid destination container and to generally obstruct the passage of fluid from the fluid destination container to the intermediate container; and

(b) instructing a user to attach the fluid transfer module to the electronically controlled fluid dispensing system to assist in transferring fluid;

wherein an interaction portion of the fluid transfer module comprises a generally transparent housing portion and a

generally opaque, generally rigid valve portion, the interaction portion being connected to the fluid transfer module such that the interaction portion is configured to permit the electronically controlled fluid dispensing system to detect that the second aperture is open when the valve portion moves into a light path in the electronically controlled fluid dispensing system; and

wherein the fluid transfer module is configured to substantially prevent fluid within the fluid transfer module from escaping upon disconnection of the fluid transfer module from the fluid source container and the fluid destination container.

2. The method of claim 1, further comprising providing the electronically controlled fluid dispensing system.

3. The method of claim 1, further comprising providing the fluid source container.

4. The method of claim 3, wherein the fluid source container is a medicinal vial.

5. The method of claim 1, further comprising providing a vial adapter configured to removably connect to the first interface between the fluid transfer module and the fluid source container, wherein the vial adapter comprises a selectively openable and closeable fluid passage that substantially prevents fluid within the vial adapter from escaping after the vial adapter is detached from the first interface.

6. The method of claim 1, further comprising providing the fluid destination container.

7. The method of claim 6, wherein the fluid destination container is an IV bag.

8. The method of claim 7, further comprising providing a selectively closeable fluid connector configured to removably connect to the second interface between the fluid transfer module and the IV bag, wherein the selectively closeable fluid connector substantially prevents fluid within the IV bag from escaping after the IV bag is detached from the second interface.

9. The method of claim 2, wherein the electronically controlled fluid dispensing system comprises a beam emitter and a beam detector, and wherein the electronically controlled fluid dispensing system detects that the second aperture is open by detecting that a beam emitted from the beam emitter has not been detected by the beam detector.

10. The method of claim 1, wherein a second interaction portion of the fluid transfer module is configured to permit the electronically controlled fluid dispensing system to detect that at least a portion of the fluid transfer module is attached to the electronically controlled fluid dispensing system, and wherein the second interaction portion of the fluid transfer module comprises a generally opaque outer housing.

11. The method of claim 1, wherein the interaction portion of the fluid transfer module comprises a closeable male luer connector.

12. The method of claim 1, wherein the fluid transfer module is configured to be disposable.

13. The method of claim 1, further comprising connecting the first interface to the fluid source container, or further comprising connecting the second interface to the fluid destination container.

14. The method of claim 2, wherein the electronically controlled fluid dispensing system is capable of detecting air bubbles in the fluid transfer module.

15. The method of claim 1, wherein the first interface is a closeable male luer connector, or wherein the second interface is a closable male luer connector.

16. The method of claim 1, wherein the fluid transfer module is configured to transfer chemotherapy fluids.

17. The method of claim 1, further comprising providing the intermediate container configured to connect to the intermediate interface.

18. The method of claim 1, wherein the fluid transfer module comprises the intermediate container.

5

19. The method of claim 1, wherein the first valve is integrally formed with the second valve as a single unitary component.

20. The method of claim 1, wherein the second interface comprises a valve member movable between a closed position and an open position, the valve member configured to be in the open position when the fluid destination container is attached to the second interface, and the valve member configured to close automatically when the fluid destination container is detached from the second interface.

10

15

21. The method of claim 2, wherein the light path in the electronically controlled fluid dispensing system is an infrared light path.

* * * * *